=> file req FILE 'REGISTRY' ENTERED AT 11:31:38 ON 18 AUG 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 American Chemical Society (ACS) => display history full 11-FILE 'HCAPLUS' ENTERED AT 10:54:29 ON 18 AUG 2006 L18130 SEA KANO ?/AU L226503 SEA ITOH ?/AU L3 2277 SEA KITAOKA ?/AU L4174 SEA YODO ?/AU L530132 SEA SHIBATA ?/AU L6 2 SEA L1 AND L2 AND L3 AND L4 AND L5 FILE 'HCA' ENTERED AT 10:56:44 ON 18 AUG 2006 L7 104989 SEA SIDECHAIN? OR SIDE#(2A)CHAIN? OR LARIAT? OR PENDANT? L8124494 SEA GRAFT? FILE 'HCAPLUS' ENTERED AT 10:57:17 ON 18 AUG 2006 SEL L6 1-2 RN FILE 'REGISTRY' ENTERED AT 10:57:39 ON 18 AUG 2006 L9 16 SEA (24980-41-4/BI OR 25248-42-4/BI OR 25322-69-4/BI OR L109 SEA L9 AND PMS/CI L11 7 SEA L9 NOT L10

FILE 'HCA' ENTERED AT 11:17:45 ON 18 AUG 2006 L12 100529 SEA L10 L13 626379 SEA POLYETHER# OR POLYESTER# OR POLY(A) (ETHER# OR ESTER#) OR POLYACRYLIC# OR POLYMETHACRYLIC# OR POLYACRYLA TE# OR POLYMETHACRYLATE# OR (ACRYLIC# OR METHACRYLIC# OR ACRYLATE# OR METHACRYLATE#) (A) (POLY OR POLYM? OR COPOLYM? OR HOMOPOLYM? OR TERPOLYM? OR RESIN? OR GUM#) L14 37939 SEA (L12 OR L13) AND (L7 OR L8) L1523245 SEA ?CARBODIIMID? L16 228 SEA L14 AND L15 L17 213162 SEA PIGMENT?

FILE 'LCA' ENTERED AT 11:19:08 ON 18 AUG 2006

L18 3311 SEA (COLOR? OR COLOUR? OR PIGMENT? OR DYE? OR STAIN? OR PAINT? OR CHROMA# OR CHROMOGEN? OR CHROMOPHOR? OR TINCT? OR TINT?)/BI,AB

FILE 'HCA' ENTERED AT 11:19:20 ON 18 AUG 2006 L19 95324 SEA CARBONBLACK# OR (CARBON# OR C) (2A) BLACK# L20 10 SEA L16 AND L17 24 SEA L16 AND L18 L21 L22 10 SEA L16 AND L19 FILE 'REGISTRY' ENTERED AT 11:22:47 ON 18 AUG 2006 E CARBON/CN L23 1 SEA CARBON/CN E GRAPHITE/CN L24 1 SEA GRAPHITE/CN FILE 'LCA' ENTERED AT 11:23:06 ON 18 AUG 2006 L25 1588 SEA L23 OR L24 OR (ACT# OR ACTIV?)(2A)(CARBON# OR C) OR GRAPHIT? OR CHARCOAL? OR CARBONIFEROUS? OR CARBONACEOUS? OR BLACKLEAD# OR LAMPBLACK# OR GASBLACK# OR CHANNELBLACK# OR STOVEBLACK# OR CHIMNEYBLACK# OR BONEBLACK# OR ANIMALBLACK# OR AMORPH? (2A) (CARBON# OR C) L26 7 SEA BLACK#(2A)(LEAD# OR LAMP# OR GAS## OR CHANNEL# OR STOVE# OR CHIMNEY# OR BONE# OR ANIMAL#) FILE 'HCA' ENTERED AT 11:28:36 ON 18 AUG 2006 L27 682750 SEA L25 OR L26 L28 5 SEA L16 AND L27 L29 33 SEA L20 OR L21 OR L22 OR L28 L30 24 SEA L29 AND 1840-2002/PY, PRY

=> file hca

FILE 'HCA' ENTERED AT 11:32:00 ON 18 AUG 2006
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=> d 130 1-24 cbib abs hitstr hitind

- L30 ANSWER 1 OF 24 HCA COPYRIGHT 2006 ACS on STN
- 141:164950 Light-shielding photosensitive resin composition and its cured product. Ito, Kazunori; Kitaoka, Naoyuki; Nabeta, Tomohiro; Hirasawa, Tamano (Sakata Inx Corp., Japan; Toyo Gosei Co., Ltd.). Jpn. Kokai Tokkyo Koho JP 2004219978 A2 20040805, 32 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 2003-280467 20030725. PRIORITY: JP 2002-379329 20021227.
- AB Title compn., which is useful in providing black matrix for **color** filter, comprises a photosensitive resin and titanium black which has

been surface-treated with a **carbodiimide** compd. having a **carbodiimide** equiv. of 100-50,000.

IT **25322-69-4DP**, Polypropylene glycol, **polycarbodiimide** -polyester-polyethers

(compn. contg. carbodiimide compd.-treated titanium black for black matrix of color filter)

RN 25322-69-4 HCA

CN Poly[oxy(methyl-1,2-ethanediyl)], α -hydro- ω -hydroxy-(9CI) (CA INDEX NAME)

$$HO = \left[(C3H6) - O \right] = H$$

IC ICM G03F007-004

ICS C08K009-00; C08L101-00; G02B005-20

CC 74-13 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)
Section cross-reference(s): 37

- ST light shielding photosensitive compn titanium black carbodiimide compd treatment
- IT Optical filters

(compn. contg. carbodiimide compd.-treated titanium black for black matrix of color filter)

IT Polyoxyalkylenes, preparation

(compn. contg. carbodiimide compd.-treated titanium black for black matrix of color filter)

IT Polyesters, preparation

(polycarbodiimide-, block; compn. contg.
carbodiimide compd.-treated titanium black for black
matrix of color filter)

IT Polyesters, preparation

(polycarbodiimide-, graft; compn. contg.
carbodiimide compd.-treated titanium black for black
matrix of color filter)

IT Acrylic polymers, preparation

(polycarbodiimide-polyesters-acrylic
polymers; compn. contg. carbodiimide
compd.-treated titanium black for black matrix of color
filter)

IT Polycarbodiimides

(polyester-, block; compn. contg. carbodiimide
compd.-treated titanium black for black matrix of color
filter)

- ITPolycarbodiimides (polyester-, graft; compn. contq. carbodiimide compd.-treated titanium black for black matrix of color filter) IT Polyoxyalkylenes, preparation (polyester-polyimide-; compn. contq. carbodiimide compd.-treated titanium black for black matrix of color filter) Polyimides, preparation ΙT (polyester-polyoxyalkylene-; compn. contq. carbodiimide compd.-treated titanium black for black matrix of color filter) ΙT Polyesters, preparation (polyimide-polyoxyalkylene-; compn. contg. carbodiimide compd.-treated titanium black for black matrix of color filter) ΙT 124-04-9DP, Adipic acid, polycarbodiimide-502-44-3DP, ε -Caprolactone, polyesters graft polycarbodiimide-polyesters polyesters 25322-69-4DP, Polypropylene glycol,
- 4457-71-0DP, 3-Methyl-1,5-pentanediol, polycarbodiimidepolycarbodiimide-polyester-polyethers (compn. contg. carbodiimide compd.-treated titanium

black for black matrix of color filter)

IT25086-15-1, Methacrylic acid-methyl methacrylate copolymer 65697-21-4, Benzyl methacrylate-methacrylic acid 67653-78-5, Dipentaerythritol hexaacrylate homopolymer 147076-20-8, Dipentaerythritol hexaacrylate-pentaerythritol triacrylate copolymer 290815-18-8, HP 100 (photoresist) 731016-17-4 731016-18-5

(compn. contg. carbodiimide compd.-treated titanium black for black matrix of color filter)

- IT 7631-86-9, Silica, uses 122026-93-1, Titanium black 13R (compn. contq. carbodimide compd.-treated titanium black for black matrix of color filter)
- L30 ANSWER 2 OF 24 HCA COPYRIGHT 2006 ACS on STN
- 141:158588 Treated titanium black, treated titanium black dispersions, and chemical compounds for titanium black treatment. Ito, Kazunori; Kitaoka, Naoyuki; Nabeta, Tomohiro (Sakata Inx Corp., Japan). Kokai Tokkyo Koho JP 2004217885 A2 20040805, 27 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 2003-278647 20030723. PRIORITY: JP 2002-379329 20021227.
- AΒ The present invention relates to treated titanium black obtained by treating the titanium black surfaces with carbodiimide compds. with ≥1 carbodiimide group and carbodiimide equiv. 100-50,000. Thus, 316 parts isocyanate group-contg. polycarbodiimide with carbodiimide

equiv., 115.7 parts poly(3-methylpentyladipate) diol 115.7, and 16 mg tetra-Bu titanate were heated at 100° for 5 h, 84.6 parts polycaprolactone having terminal carboxylic acid groups was added therein and reacted at 80° for 2 h, 375.5 parts propylene glycol monomethyl ether acetate was added therein to give a 40%-solids carbodiimide compd. with no. av. mol. wt. 4200 and carbodiimide equiv. 1583, 50 g amorphous silica-coated Titanium Black 13R was dispersed in 450 g water, 15 g of the resulting carbodiimide compd. soln. and 170 g propylene glycol monomethyl ether acetate were added therein and reacted at 90° for 8 h, dried at 80° for 2 h to give surface-treated titanium black, 48.0 parts of which was mixed with 52.0 parts propylene glycol monomethyl ether acetate, 35.0 parts the resulting dispersion was mixed with 30.0 parts a polyurethane varnish and 35.0 parts a mixed solvent, showing good dispersion stability and elec. resistance ≥1013 for a printing ink film.

IT 24980-41-4DP, Polycaprolactone, carboxy-terminated, reaction products with isocyanate-contg. polycarbodiimide and polyester diols 39751-34-3DP, Adipic

acid-3-methyl-1,5-pentanediol copolymer, diols, reaction products with isocyanate-contg. **polycarbodiimide** and polycaprolactone

(assumed monomers; prepn. of **carbodiimide** surface-treated titanium black)

RN 24980-41-4 HCA

CN 2-Oxepanone, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 502-44-3 CMF C6 H10 O2

RN 39751-34-3 HCA

CN Hexanedioic acid, polymer with 3-methyl-1,5-pentanediol (9CI) (CA INDEX NAME)

CM 1

CRN 4457-71-0 CMF C6 H14 O2

$$^{\rm Me}_{\rm HO-CH_2-CH_2-CH_2-CH_2-OH}$$

CM 2

CRN 124-04-9 CMF C6 H10 O4

 $HO_2C - (CH_2)_4 - CO_2H$

25248-42-4DP, Polycaprolactone, carboxy-terminated, reaction products with isocyanate-contg. polycarbodiimide and polyester diols 25322-69-4DP, Polypropylene glycol, carboxy-terminated, reaction products with polycarbodiimdes and polyester diols 58991-77-8DP, Adipic acid-3-methyl-1,5-pentanediol copolymer, sru, diols, reaction products with isocyanate-contg. polycarbodiimide and polycaprolactone

(prepn. of **carbodiimide** surface-treated titanium black) 25248-42-4 HCA

CN Poly[oxy(1-oxo-1,6-hexanediyl)] (9CI) (CA INDEX NAME)

RN 25322-69-4 HCA

RN

CN Poly[oxy(methyl-1,2-ethanediyl)], α -hydro- ω -hydroxy-(9CI) (CA INDEX NAME)

$$HO \longrightarrow (C3H6) - O \longrightarrow D$$

RN 58991-77-8 HCA

```
CN
    Poly[oxy(3-methyl-1,5-pentanediyl)oxy(1,6-dioxo-1,6-hexanediyl)]
     (9CI) (CA INDEX NAME)
  IC
    ICM C09C001-36
     ICS C09C003-06; C09C003-10; C09C003-12
CC
     42-6 (Coatings, Inks, and Related Products)
     Section cross-reference(s): 74, 76
    treated titanium black dispersion chem compd treatment;
ST
    polyester polycarbodiimide coated titanium black
    dispersion ink compn
IT
    Polyesters, uses
        (acrylic, polycarbodiimide-; prepn. of
       carbodiimide surface-treated titanium black)
ΙT
    Acrylic polymers, uses
        (carboxy-terminated, reaction products with polycarbodiimdes and
       polyester diols; prepn. of carbodiimide
       surface-treated titanium black)
TT
     Silanes
        (elec. insulators, titanium black coated with; prepn. of
       carbodiimide surface-treated titanium black)
ΙT
    Acrylic polymers, uses
        (graft, polycarbodiimide-; prepn. of
       carbodiimide surface-treated titanium black)
IT
     Binders
        (ink compns.; prepn. of carbodiimide surface-treated
       titanium black)
ΙT
     Polyesters, uses
      Polyethers, uses
        (polycarbodiimide-, graft; prepn. of
       carbodiimide surface-treated titanium black)
ΙT
    Polyesters, uses
        (polycarbodiimide-; prepn. of carbodiimide
        surface-treated titanium black)
IT
     Polyurethanes, uses
        (polyester-, binder for inks; prepn. of
       carbodiimide surface-treated titanium black)
ΙT
     Polycarbodiimides
        (polyester-, graft; prepn. of
       carbodiimide surface-treated titanium black)
IT
    Acrylic polymers, uses
     Polyoxyalkylenes, uses
```

```
(polyester-, polycarbodiimide-; prepn. of
        carbodiimide surface-treated titanium black)
IT
     Polycarbodiimides
        (polyether-, graft; prepn. of
        carbodiimide surface-treated titanium black)
ΙT
     Polyesters, uses
        (polyoxyalkylene-, polycarbodiimide-; prepn. of
        carbodiimide surface-treated titanium black)
     Dispersion (of materials)
ΙT
     Glass substrates
     Resists
     Surface treatment
        (prepn. of carbodiimide surface-treated titanium black)
IT
     Polyoxyalkylenes, uses
        (prepn. of carbodimide surface-treated titanium black)
IΤ
     Inks
        (printing; prepn. of carbodiimide surface-treated
        titanium black)
ΙT
     Polycarbodiimides
        (reaction products with polyester diols and
        carboxy-terminated polycaprolactone; prepn. of
        carbodiimide surface-treated titanium black)
ΙT
     Electric insulators
        (titanium black coated with; prepn. of carbodismide
        surface-treated titanium black)
ΙT
     Pigments, nonbiological
        (treated; prepn. of carbodiimide surface-treated
        titanium black)
ΙT
     7631-86-9, Amorphous silica, uses
        (amorphous, elec. insulator, titanium black coated with; prepn.
        of carbodiimide surface-treated titanium black)
ΙT
     24980-41-4DP, Polycaprolactone, carboxy-terminated, reaction
     products with isocyanate-contg. polycarbodiimide and
     polyester diols 39751-34-3DP, Adipic
     acid-3-methyl-1,5-pentanediol copolymer, diols, reaction products
     with isocyanate-contg. polycarbodiimide and
     polycaprolactone
        (assumed monomers; prepn. of carbodiimide
        surface-treated titanium black)
ΙT
     136020-57-0P, Adipic acid-isophoronediamine-isophorone
     diisocyanate-neopentyl glycol copolymer
        (composed of actual and assumed monomers, binder for ink; prepn.
        of carbodiimide surface-treated titanium black)
IT
     122026-93-1, Titanium Black 13R
        (pigment, surface-treated; prepn. of
        carbodiimide surface-treated titanium black)
ΙT
     112540-76-8, Titanium black
```

(pigment; prepn. of carbodiimide
surface-treated titanium black)

105-59-9DP, Methyldiethanolamine, reaction products with polycarbodiimde and polyester diols 25248-42-4DP, Polycaprolactone, carboxy-terminated, reaction products with isocyanate-contg. polycarbodiimide and polyester diols 25322-69-4DP, Polypropylene glycol, carboxy-terminated, reaction products with polycarbodiimdes and polyester diols 58991-77-8DP, Adipic acid-3-methyl-1,5-pentanediol copolymer, sru, diols, reaction products with isocyanate-contg. polycarbodiimide and polycaprolactone

(prepn. of carbodiimide surface-treated titanium black)

- IT 67653-78-5P, Dipentaerythritol hexaacrylate homopolymer (resist binder; prepn. of **carbodiimide** surface-treated titanium black)
- IT 65697-21-4, Benzyl methacrylate-methacrylic acid copolymer (resist binder; prepn. of **carbodiimide** surface-treated titanium black)
- IT 9003-07-0, P 2161 (substrate; prepn. of **carbodiimide** surface-treated titanium black)
- L30 ANSWER 3 OF 24 HCA COPYRIGHT 2006 ACS on STN
- 140:289965 Adhesive film for sealing metal terminal for lithium battery. Mochizuki, Yoichi; Okushita, Masataka (Dainippon Printing Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 2004095543 A2 20040325, 20 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 2003-283295 20030731. PRIORITY: JP 2002-228430 20020806.
- AB The sealing film for metal electrode terminals protruding from a laminated battery package, contg. an inner hot adhering polyolefin resin layer and a metal foil barrier layer, is a biaxial extended poly(ethylene naphthalate) film covered with polyolefin on both sides, with the polyolefin at least on 1 side being an acid modified polyolefin.
- IC ICM H01M002-06 ICS H01M002-08; H01M010-40; C09J007-00; C09J123-26
- CC 52-1 (Electrochemical, Radiational, and Thermal Energy Technology)
- IT Epoxy resins, uses

Polyesters, uses

(adhesion promotor; adhesive film for sealing metal terminal for lithium battery)

IT Carbon black, uses

(adhesive film for sealing metal terminal for lithium battery)

IT Acrylic polymers, uses

(amino group **grafted**, adhesion promotor; adhesive film for sealing metal terminal for lithium battery)

```
2422-91-5, Triphenylmethane-4,4',4''-triisocyanate
IΤ
     Tris(p-isocyanatophenyl)thiophosphate 7723-14-0D, Phosphorus,
               9002-98-6
                          9016-87-9, Polymethylenepolyphenylpolyisocyana
          16065-83-1D, Chromium (III), compds., uses
                                                      25215-75-2,
     Polycarbodiimide
                        27598-85-2, Aminophenol
        (adhesion promotor; adhesive film for sealing metal terminal for
        lithium battery)
L30
    ANSWER 4 OF 24 HCA COPYRIGHT 2006 ACS on STN
140:43588 Treated pigment, use thereof, and compounds for
    pigment treatment. Kano, Masanori; Itoh, Kazunori; Yodo,
     Takaaki; Kitaoka, Naoyuki; Shibata, Hideo (Sakata Inx Corp., Japan).
       PCT Int. Appl. WO 2004000950 A1 20031231, 61 pp.
                                                        DESIGNATED
     STATES: W: CN, JP, KR, US; RW: AT, BE, CH, CY, DE, DK, ES, FI, FR,
     GB, GR, IE, IT, LU, MC, NL, PT, SE, TR. (Japanese). CODEN: PIXXD2.
     APPLICATION: WO 2003-JP8019 20030625. PRIORITY: JP 2002-185172
     20020625.
AB
     A treated pigment having satisfactory dispersibility in a dispersion
     medium and flowability can be obtained even from a pigment having, on
     the surface, no functional group reactive with a carbodiimide group.
     The treated pigment is used in a dispersion compn. and also in a
     resist compn. which has excellent developing characteristics when
     used in forming a pattern, e.g., a color filter or black matrix,
     therefrom. Also provided is a compd. for pigment treatment which is
     suitable for producing the treated pigment. The treated pigment is
     obtained by treating a pigment with a carbodiimide compd. having ≥1
     side chains (e.g., a polyester side chain, polyether side chain,
     polyacrylic side chain) and ≥1 basic nitrogen-contg. groups.
ΙT
     9003-11-6DP, Ethylene oxide-propylene oxide copolymer,
     carboxy group-terminated, reaction products with
    polycarbodiimide derivs. 9011-14-7DP, Poly(methyl
    methacrylate), diol derivs. or carboxyl group-terminated,
     graft product with polycarbodiimides derivs.
     24980-41-4DP, Polycaprolactone, graft products
     with polycarbodimide derivs. 25248-42-4DP,
     Polycaprolactone, graft products with
    polycarbodiimide derivs.
        (compd. for pigment treatment; treated pigment
        for dispersion compn. and resist compn.)
RN
     9003-11-6 HCA
CN
    Oxirane, methyl-, polymer with oxirane (9CI) (CA INDEX NAME)
     CM
         1
     CRN 75-56-9
```

CMF C3 H6 O

CM 2

CRN 75-21-8 CMF C2 H4 O

 $\overset{\circ}{\sim}$

RN 9011-14-7 HCA

CN 2-Propenoic acid, 2-methyl-, methyl ester, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 80-62-6 CMF C5 H8 O2

RN 24980-41-4 HCA

CN 2-Oxepanone, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 502-44-3 CMF C6 H10 O2

```
RN
    25248-42-4 HCA
CN
    Poly[oxy(1-oxo-1,6-hexanediyl)] (9CI) (CA INDEX NAME)
IC
    ICM C09C003-08
     ICS C09C003-10
CC
    42-6 (Coatings, Inks, and Related Products)
     Section cross-reference(s): 74
ST
    pigment dispersibility flowability improvement
    polycarbodiimide surface treatment
ΙT
    Carbon black, uses
        (pigment; treated pigment for dispersion
       compn. and resist compn.)
IT
    Polyesters, uses
      Polyethers, uses
        (polycarbodiimide-, graft, for
       pigment treatment; treated pigment for
       dispersion compn. and resist compn.)
ΙT
    Polycarbodiimides
        (polyester-, graft, for pigment
       treatment; treated pigment for dispersion compn. and
        resist compn.)
IT
    Polycarbodiimides
        (polyether-, graft, for pigment
       treatment; treated pigment for dispersion compn. and
        resist compn.)
    Optical filters
IT
     Photoresists
      Pigments, nonbiological
        (treated pigment for dispersion compn. and resist
       compn.)
ΙT
     103-74-2DP, 2-Pyridineethanol, reaction products with
    polycarbodiimide graft copolymers 105-59-9DP,
    Methyldiethanolamine, reaction products with
    polycarbodiimide graft copolymers
                                        108-30-5DP,
    Succinic anhydride, reaction products with polycarbodiimide
                       4098-71-9DP, IPDI, reaction products with
     graft copolymers
    polycarbodiimide derivs. 9003-11-6DP, Ethylene
    oxide-propylene oxide copolymer, carboxy group-terminated, reaction
    products with polycarbodiimide derivs. 9011-14-7DP
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, Poly(methyl methacrylate), diol derivs. or carboxyl group-terminated, graft product with polycarbodiimides derivs. 24980-41-4DP, Polycaprolactone, graft products with polycarbodiimide derivs. 25248-42-4DP, Polycaprolactone, graft products with polycarbodiimide derivs.

(compd. for **pigment** treatment; treated **pigment** for dispersion compn. and resist compn.)

- L30 ANSWER 5 OF 24 HCA COPYRIGHT 2006 ACS on STN
 139:262267 Treated pigments with good dispersibility, their
 uses and the treating compounds therefor. Kano, Masanori; Itoh,
 Kazunori; Kitaoka, Naoyuki; Yodo, Takaaki; Shibata, Hideo (Sakata
 Inx Corp., Japan). PCT Int. Appl. WO 2003076527 A1 20030918, 57 pp.
 DESIGNATED STATES: W: CN, JP, KR, US; RW: AT, BE, CH, CY, DE, DK,
 ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR. (Japanese).
 CODEN: PIXXD2. APPLICATION: WO 2003-JP2772 20030310. PRIORITY: JP
 2002-64416 20020308; JP 2002-74966 20020318; JP 2002-89231 20020327;
 JP 2002-186838 20020626.
- Title pigments are carbodiimide-reactive functional group-contg. org. pigments or carbon black treated with carbodimides contg. polyester, polyether, and/or acrylic side chains and having carbodimido equiv. (EQ) of 100-50,000. An aq. carbon black (with pH 3.1) dispersion was stirred with a glycol ether acetate and a compd. [with EQ 2,053; prepd. from NCO-contg. polycarbodiimide, poly(3-methylpentyl adipate) diol, and polycaprolactone] at 90° for 8 h and vacuum dried to form a dispersion showing no ppt. and no viscosity increase after 1 wk at room temp., which was used to prep. a resist with high optical d., resistivity, image development.
- 9003-63-8DP, Poly(butyl methacrylate), diol derivs., polymers with NCO-contg. polycarbodiimides and polycaprolactones 24980-41-4DP, polymers with NCO-contg. polycarbodiimides and poly(methylpentyl adipate) diols or polyols 25248-42-4DP, Polycaprolactone, polymers with NCO-contg. polycarbodiimides and poly(methylpentyl adipate) diols or polyols 25322-69-4DP, Polypropylene glycol, COOH-terminated, polymers with NCO-contg. polycarbodiimides and poly(methylpentyl adipate) diols 58991-77-8DP, Adipic acid-3-methyl-1,5-pentanediol copolymer, sru, diol derivs., polymers with NCO-contg. polycarbodiimides and polycaprolactones

(acrylic, polyester, and/or polyether
side chain-contg. carbodiimide
-treated functional pigments or carbon
black for dispersion stability)
9003-63-8 HCA

CN 2-Propenoic acid, 2-methyl-, butyl ester, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 97-88-1 CMF C8 H14 O2

RN 24980-41-4 HCA CN 2-Oxepanone, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 502-44-3 CMF C6 H10 O2



RN 25248-42-4 HCA

CN Poly[oxy(1-oxo-1,6-hexanediyl)] (9CI) (CA INDEX NAME)

RN 25322-69-4 HCA

CN Poly[oxy(methyl-1,2-ethanediyl)], α -hydro- ω -hydroxy-(9CI) (CA INDEX NAME)

$$HO - (C3H6) - O - n$$

RN 58991-77-8 HCA
CN Poly[oxy(3-methyl-1,5-pentanediyl)oxy(1,6-dioxo-1,6-hexanediyl)]
(9CI) (CA INDEX NAME)

39751-34-3DP, Adipic acid-3-methyl-1,5-pentanediol copolymer, diol derivs., polymers with NCO-contg. polycarbodiimides and polycaprolactones (assumed monomers; acrylic, polyester, and/or

polyether side chain-contg.

carbodiimide-treated functional pigments or

carbon black for dispersion stability)

RN 39751-34-3 HCA

CN Hexanedioic acid, polymer with 3-methyl-1,5-pentanediol (9CI) (CA INDEX NAME)

CM 1

CRN 4457-71-0 CMF C6 H14 O2

$$Me$$
 $HO-CH_2-CH_2-CH-CH_2-CH_2-OH$

CM 2

CRN 124-04-9 CMF C6 H10 O4

 $HO_2C - (CH_2)_4 - CO_2H$

RN 196402-77-4 HCA

2-Propenoic acid, 2-methyl-, polymer with 2-[[3-[(1-oxo-2-propenyl)oxy]-2,2-bis[[(1-oxo-2-propenyl)oxy]methyl]propoxy]methyl]2-[[(1-oxo-2-propenyl)oxy]methyl]-1,3-propanediyl di-2-propenoate and phenylmethyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 29570-58-9 CMF C28 H34 O13

CM 2 ·

CRN 2495-37-6 CMF C11 H12 O2

CM 3

CRN 79-41-4 CMF C4 H6 O2

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CH2
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Me-C-CO2H
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IC
     ICM C09C003-10
     ICS C09C001-48; G03F007-004; G02B005-20; G02B005-00
     42-6 (Coatings, Inks, and Related Products)
CC
     Section cross-reference(s): 74
ST
    polyester carbodiimide treated functional
    pigment dispersion stability; polyether
     carbodiimide treated functional pigment dispersion
     stability; acrylic carbodiimide treated functional
    pigment dispersion stability; carbon black
    polyester carbodiimide treatment dispersion
     stability; image development resist polyester
    carbodiimide treated functional pigment
     Polyoxyalkylenes, uses
IT
        (COOH-terminated, polymers with NCO-contg.
        polycarbodiimides and poly(methylpentyl adipate) diols;
        acrylic, polyester, and/or polyether
        side chain-contq. carbodiimide
        -treated functional pigments or carbon
        black for dispersion stability)
IT
     Pigments, nonbiological
     Resists
        (acrylic, polyester, and/or polyether
        side chain-contg. carbodiimide
        -treated functional pigments or carbon
        black for dispersion stability)
IT
    Carbon black, uses
        (acrylic, polyester, and/or polyether
        side chain-contg. carbodiimide
        -treated functional pigments or carbon
        black for dispersion stability)
TI
     Polyoxyalkylenes, uses
        (acrylic-polycarbodiimide-; acrylic, polyester
        , and/or polyether side chain
        -contg. carbodiimide-treated functional
        pigments or carbon black for
        dispersion stability)
IT
    Polycarbodiimides
        (acrylic-polyoxyalkylene-; acrylic, polyester, and/or
        polyether side chain-contg.
        carbodiimide-treated functional pigments or
        carbon black for dispersion stability)
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IT
     Polyurethanes, uses
        (polycarbodiimide-polyester-; acrylic,
        polyester, and/or polyether side
        chain-contg. carbodiimide-treated functional
        pigments or carbon black for
        dispersion stability)
ΙT
     Polyurethanes, uses
        (polycarbodiimide-polyester-polyether
        -; acrylic, polyester, and/or polyether
        side chain-contq. carbodiimide
        -treated functional pigments or carbon
        black for dispersion stability)
ΙT
     Polyethers, uses
        (polycarbodiimide-polyester-polyurethane-;
        acrylic, polyester, and/or polyether
        side chain-contg. carbodiimide
        -treated functional pigments or carbon
        black for dispersion stability)
IT
     Polyesters, uses
        (polycarbodiimide-polyether-polyurethane-;
        acrylic, polyester, and/or polyether
        side chain-contg. carbodiimide
        -treated functional pigments or carbon
        black for dispersion stability)
IT
     Acrylic polymers, uses
        (polycarbodiimide-polyoxyalkylene-; acrylic,
        polyester, and/or polyether side
        chain-contg. carbodiimide-treated functional
        pigments or carbon black for
        dispersion stability)
ΙT
     Polyesters, uses
        (polycarbodiimide-polyurethane-; acrylic,
        polyester, and/or polyether side
        chain-contg. carbodiimide-treated functional
        pigments or carbon black for
        dispersion stability)
IT
     Polycarbodiimides
        (polyester-polyether-polyurethane-; acrylic,
        polyester, and/or polyether side
        chain-contq. carbodiimide-treated functional
        pigments or carbon black for
        dispersion stability)
IT
     Polycarbodiimides
        (polyester-polyurethane-; acrylic, polyester,
        and/or polyether side chain-contg.
        carbodiimide-treated functional pigments or
        carbon black for dispersion stability)
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ΙT
     Polyoxyalkylenes, uses
        (polymers with NCO-contg. polycarbodiimides and
       polycaprolactones; acrylic, polyester, and/or
       polyether side chain-contg.
       carbodiimide-treated functional pigments or
        carbon black for dispersion stability)
IT
     79-10-7DP, Acrylic acid, COOH-contg. polymers, polymers with
    NCO-contg. polycarbodiimides and poly(methylpentyl
    adipate) diols
                      79-41-4DP, Methacrylic acid, COOH-contq. polymers,
    polymers with NCO-contg. polycarbodiimides and
    poly(methylpentyl adipate) diols
                                        107-21-1DP, Ethylene glycol,
    polymers with NCO-contg. polycarbodiimides and
    poly(methylpentyl adipate) diols and polycaprolactones
     9003-63-8DP, Poly(butyl methacrylate), diol derivs.,
    polymers with NCO-contg. polycarbodiimides and
    polycaprolactones 24980-41-4DP, polymers with NCO-contq.
    polycarbodiimides and poly(methylpentyl adipate) diols or
    polyols 25248-42-4DP, Polycaprolactone, polymers with
    NCO-contg. polycarbodiimides and poly(methylpentyl
    adipate) diols or polyols 25322-69-4DP, Polypropylene
    glycol, COOH-terminated, polymers with NCO-contg.
    polycarbodiimides and poly(methylpentyl adipate) diols
    25322-69-4DP, Polypropylene glycol, polymers with NCO-contg.
    polycarbodiimides and polycaprolactones 58991-77-8DP
     , Adipic acid-3-methyl-1,5-pentanediol copolymer, sru, diol derivs.,
    polymers with NCO-contg. polycarbodiimides and
    polycaprolactones
        (acrylic, polyester, and/or polyether
        side chain-contg. carbodiimide
        -treated functional pigments or carbon
       black for dispersion stability)
ΙT
    39751-34-3DP, Adipic acid-3-methyl-1,5-pentanediol
    copolymer, diol derivs., polymers with NCO-contg.
    polycarbodiimides and polycaprolactones
        (assumed monomers; acrylic, polyester, and/or
       polyether side chain-contq.
       carbodiimide-treated functional pigments or
        carbon black for dispersion stability)
ΙT
    196402-77-4, Benzyl methacrylate-dipentaerythritol
    hexaacrylate-methacrylic acid copolymer
        (resist binder; acrylic, polyester, and/or
       polyether side chain-contg.
       carbodiimide-treated functional pigments or
       carbon black for dispersion stability)
    ANSWER 6 OF 24 HCA COPYRIGHT 2006 ACS on STN
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138:137713 Spatially resolved derivatization of solid-phase-synthesis

beads with fluorescent dendrimers: creation of localized microdomains. Cardona, Claudia M.; Jannach, Stephan H.; Huang, Hao; Itojima, Yukiko; Leblanc, Roger M.; Gawley, Robert E.; Baker, Gary A.; Brauns, Eric B. (Department of Chemistry, Freshwater Biomedical Sciences Center, University of Miami, Coral Gables, FL, 33124-0431, USA). Helvetica Chimica Acta, 85(10), 3532-3558 (English) 2002. CODEN: HCACAV. ISSN: 0018-019X. Publisher: Verlag Helvetica Chimica Acta.

- AB Second and third generation Newkome-type trifurcated dendrimers, contg. either a coumarin or dansyl fluorescent probe at the dendrimer core, have been synthesized and attached to ArgoGel solid-phase-synthesis beads. Subsequent reaction with rhodamine dye shows that the dye can penetrate throughout the beads to acylate the remaining sites. Thus, it is possible to achieve a spatially resolved microdomain for library formation at the core of the dendrimer, primarily on the bead's periphery, and a second microdomain suitable for derivatization by other reagents such as encoding tags and fluorescent sensors.
- CC 35-8 (Chemistry of Synthetic High Polymers)
 Section cross-reference(s): 38
- ST derivatization solid phase bead fluorescent dendrimer; coumarin contg dendrimer solid phase bead; dansyl contg dendrimer solid phase bead; oxirane styrene **graft** copolymer bead fluorescent dendrimer
- IT Fluorescence

(of coumarin- and dansyl-contg. fluorescent **dye**Newkome-type trifurcated dendrimer-attached ArgoGel beads)

IT Polyethers, preparation

(polyamide-, dendrimers; prepn. of coumarin- and dansyl-contg. fluorescent **dye** Newkome-type trifurcated dendrimer-attached ArgoGel beads)

IT Dendritic polymers

(polyamide-polyethers; prepn. of coumarin- and dansyl-contg. fluorescent dye Newkome-type trifurcated dendrimer-attached ArgoGel beads)

IT Polyamides, preparation

(polyether-, dendrimers; prepn. of coumarin- and dansyl-contg. fluorescent dye Newkome-type trifurcated dendrimer-attached ArgoGel beads)

IT Polyoxyalkylenes, preparation

(polystyrene-, graft, reaction products with coumarin- or dansyl-contg. dendrimers; prepn. and characterization of)

IT 493006-92-1P

(fluorescent dyes; prepn. and characterization of)

IT 493007-16-2P

(fluorescent dyes; prepn. and characterization of)

IT 81-88-9DP, Rhodamine B, reaction products with coumarin- and

dansyl-contg. fluorescent **dye**-attached ArgoGel beads 117381-20-1DP, ArgoGel, amino-contg., reaction products with coumarin- or dansyl-contg. fluorescent **dye** Newkome-type trifurcated dendrimers 493006-92-1DP, reaction products with ArgoGel 493007-16-2DP, reaction products with ArgoGel 493007-24-2P 493007-30-0P 493007-33-3P

IT 200133-18-2P

(prepn. and reaction with coumarin-contg. fluorescent **dye** and with diisopropyl **carbodiimide**)

- IT 693-13-0, Diisopropyl carbodiimide (reaction with nonacid deriv.)
- L30 ANSWER 7 OF 24 HCA COPYRIGHT 2006 ACS on STN

(prepn. and characterization of)

- 137:301741 Development of thermally stable novel EO-polymers. Ushiwata, Takami; Okamoto, Etsuya; Kaino, Toshikuni (Institute of Material Research for Advanced Materials, Tohoku University, Sendai, 980-8577, Japan). Molecular Crystals and Liquid Crystals Science and Technology, Section A: Molecular Crystals and Liquid Crystals, 374, 303-314 (English) 2002. CODEN: MCLCE9. ISSN: 1058-725X. Publisher: Taylor & Francis Ltd..
- The authors synthesized and evaluated NLO properties of azo dye AΒ attached novel NLO polymers with high thermal stability. The NLO polymers contg. an azo dye, DR-1, in their side-chain were synthesized using a new approach. Polyacrylic acid or polymethacrylic acid reacted with DR-1 via the intermediate reaction of the acid and N, N'- dicyclohexylcarbodiimide. The DR-1 contents of the products were 39 mol% and 29 mol%, for derivs. from polyacrylic acid and polymethacrylic acid, resp. Glass transition temps. of 150° to 170° were obtained for Disperse red 1 dye attached polymethacrylic acid. The poled film exhibited 2nd order NLO susceptibilities ($\chi(2)$ 33) of 48 pm/V to 53 pm/V at 1.3 μ m fundamental wave. Due to the high glass transition temps. of the polymers, long-term stability of the optical nonlinearity at 100° was obsd. for 200 h or more. However residual carboxyl groups caused absorbance decrease mainly by hydrolysis of the ester bonds of the polymers. These polymers can be further improved to have more thermally stable structure by heat treatment. Through the approach, methacryl imide structure was formed and the polymer has high Tg of 165° . This imidized polymer exhibited $\chi(2)33$ of 45 pm/V at a wavelength of 1.3 μ m and maintained .apprx.90% of the initial value after 230 h or more at 100°.
- CC 73-10 (Optical, Electron, and Mass Spectroscopy and Other Related Properties)
- ST thermal stability nonlinear optical polymer disperse red azo dye; imidization UV visible absorption
- IT Azo **dyes**Imidation

Nonlinear optical susceptibility
Thermal stability
UV and visible spectra
 (development of thermally stable novel nonlinear optical polymers)

L30 ANSWER 8 OF 24 HCA COPYRIGHT 2006 ACS on STN 137:59847 Active and biocompatible platforms prepa

137:59847 Active and biocompatible platforms prepared by polymerization of surface coating films. Huang, Mingxian; Wang, Xiaobo; Wu, Lei; Yang, Weiping; Cheng, Jing (Aviva Biosciences, USA). PCT Int. Appl. WO 2002052045 Al 20020704, 82 pp. DESIGNATED STATES: W:
AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG, TR. (English). CODEN: PIXXD2.

NE, NL, PT, SE, SN, TD, TG, TR. (English). CODEN: PIXXD2. APPLICATION: WO 2001-US48919 20011213. PRIORITY: US 2000-258281P 20001226.

The present invention recognizes that polymerizable coating films can be utilized to make chips such as biochips that include channel structures. These chips can optionally include one or more addnl. structures such as particles, biol. groups or chem. groups. Such biochips having channel structures have a wide variety of useful applications, particularly in the field of lab. on a chip and other applications where microfluidics are of importance. One aspect of the present invention is a platform that includes: a surface, a coating film and a channel structure. Preferably, the coating film defines in part said channel structure and more preferably the platform comprises a microchip. Diagrams describing the app. assembly and operation are given.

TT 7440-44-0, Carbon, uses 9011-14-7, PMMA
 (active and biocompatible platforms prepd. by polymn.
 of surface coating films)

RN 7440-44-0 HCA

CN Carbon (7CI, 8CI, 9CI) (CA INDEX NAME)

С

RN 9011-14-7 HCA

CN 2-Propenoic acid, 2-methyl-, methyl ester, homopolymer (9CI) (CA INDEX NAME)

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CM
          1
     CRN 80-62-6
     CMF C5 H8 O2
  H2C O
IC
     ICM C12Q001-68
CC
     9-1 (Biochemical Methods)
     Section cross-reference(s): 36
IT
     Acrylic polymers, uses
     Fluoropolymers, uses
     Glass, uses
     Metals, uses
     Oxides (inorganic), uses
     Plastics, uses
     Polyimides, uses
     Polyoxyalkylenes, uses
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Polysaccharides, uses Polysiloxanes, uses

(polymers of $pendant \ \alpha.\beta$ unsatd.; active and biocompatible platforms prepd. by polymn. of surface coating films)

9003-06-9, Poly(acrylamide/acrylic acid) 25952-53-8, 1-Ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride 72607-53-5, N-(3-Aminopropyl)methacrylamide hydrochloride 82436-77-9, Bis(sulfosuccinimidyl)suberate (active and biocompatible platforms prepd. by polymn. of surface coating films)

ANSWER 9 OF 24 HCA COPYRIGHT 2006 ACS on STN L30 136:63287 A novel contamination sensor in solution: the response of the electric resistance of a composite based on crystalline polymergrafted carbon black. Tsubokawa, N.; Tsuchida, M.; Chen, J.; Nakazawa, Y. (Department of Material Science and Technology, Faculty of Engineering, Niigata University, Niigata, 950-2181, Japan). Sensors and Actuators, B: Chemical, B79(2-3), 92-97 (English) 2001. CODEN: SABCEB. ISSN: 0925-4005. Publisher: Elsevier Science B.V.. AΒ The response of the elec. resistance of the composite prepd. from cryst. polymer-grafted carbon black as a function of the contamination in soln. was studied. The elec. resistance of the composite prepd. from poly(&-caprolactone) (PCL)-grafted carbon black drastically increased in hexane, contq. chloroform, trichloroethane, and ethanol and returned immediately to the initial resistance when it was transferred into pure hexane. The response of the elec. resistance was highly reproducible. The logarithm of elec. resistance was linearly proportional to the concn. of chloroform in Also, the elec. resistance of the composite prepd. from cryst. poly(ethylene glycol) (PEG)-grafted and poly(ethylene adipate) (PEA) -grafted carbon black also drastically increased in hexane contg. alc., which is a good solvent of the grafted chain (PEG and PEA). This is due to a widening of the gaps between the carbon black particles based on the absorption of solvent for these cryst. polymers. IT24980-41-4D, Poly(ε-caprolactone), carbon black grafted with 25248-42-4D, Poly[oxy(1-oxo-1,6-hexanediyl)], carbon black grafted with (response of elec. resistance of composite based on cryst. polymer-grafted carbon black for novel contamination sensor in soln.)

CM 1

24980-41-4

HCA

2-Oxepanone, homopolymer (9CI) (CA INDEX NAME)

CRN 502-44-3 CMF C6 H10 O2

RN

CN

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RN
    25248-42-4 HCA
CN
    Poly[oxy(1-oxo-1,6-hexanediyl)] (9CI) (CA INDEX NAME)
CC
    80-2 (Organic Analytical Chemistry)
    Section cross-reference(s): 76
ST
    contamination sensor elec resistance cryst polymer grafted
    carbon black
    Polyoxyalkylenes, analysis
IT
        (carbon black grafted with;
       response of elec. resistance of composite based on cryst.
       polymer-grafted carbon black for
       novel contamination sensor in soln.)
ΙT
    Carbon black, analysis
        (cryst. polymer-grafted; response of elec. resistance
       of composite based on cryst. polymer-grafted
       carbon black for novel contamination sensor in
       soln.)
IT
    Electric resistance
    Sensors
        (response of elec. resistance of composite based on cryst.
       polymer-grafted carbon black for
       novel contamination sensor in soln.)
ΙT
    64-17-5, Ethanol, analysis
                                 67-66-3, Chloroform, analysis
    25323-89-1, Trichloroethane
        (analyte; response of elec. resistance of composite based on
       cryst. polymer-grafted carbon black
       for novel contamination sensor in soln.)
ΙT
    538-75-0, N, N'-Dicyclohexylcarbodiimide
        (condensing agent in prepn. of cryst. polymer-grafted
       carbon black)
ΙT
    24937-05-1D, Poly(ethylene adipate), carbon black
    grafted with 24938-37-2D, Poly(ethylene adipate),
    carbon black grafted with
    24980-41-4D, Poly(\epsilon-caprolactone), carbon
    black grafted with 25248-42-4D,
    Poly[oxy(1-oxo-1,6-hexanediyl)], carbon black
    grafted with 25322-68-3D, Poly(ethylene glycol),
    carbon black grafted with
        (response of elec. resistance of composite based on cryst.
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polymer-grafted carbon black for
        novel contamination sensor in soln.)
ΙT
     110-54-3, Hexane, analysis
        (sample solvent; response of elec. resistance of composite based
        on cryst. polymer-grafted carbon
        black for novel contamination sensor in soln.)
L30
    ANSWER 10 OF 24
                      HCA COPYRIGHT 2006 ACS on STN
135:212334
          Postgrafting of hydrophobic polymers to hydrophilic polymer-
     grafted carbon black and their
     dispersibility.
                      Tsubokawa, Norio; Ohmata, Kimiyo; Akai, Katsunori;
     Nishizawa, Nobuyoshi; Fujiki, Kazuhiro (Department of Materials
     Science and Technology, Faculty of Engineering, Niigata University,
     Ikagashi, Niigata, 950-2181, Japan). Shikizai Kyokaishi, 74(1), 2-7
     (Japanese) 2001. CODEN: SKYOAO. ISSN: 0010-180X.
     Publisher: Shikizai Kyokai.
AΒ
     Carbon black was grafted with poly(ethyleneimine) (I) and glycidyl
     methacrylate-Me methacrylate copolymer (II). Carbon black grafted
     with I gave a stable colloidal dispersion in water, but pptd. in
     toluene.
               Carbon black grafted with I and II lost dispersibility in
     water, but uniformly and stably dispersed in toluene.
                                                            When carbon
     black grafted with I and II was dispersed in a water-toluene binary
     mixt., the water droplets (.apprx.1-3-mm diam.) were stabilized and
     gathered in the interfacial layer. This result indicated that the
     water droplets were surrounded by the particles in the same manner as
     surfactant mols.
CC
     42-6 (Coatings, Inks, and Related Products)
ST
     carbon black ethyleneimine glycidyl methacrylate
     grafting; hydrophilicity hydrophobicity grafted
     carbon black
ΙT
    Carbon black, properties
        (FW 200, reaction products with glycidyl methacrylate-Me
       methacrylate copolymer and poly(ethyleneimine);
        postgrafting of hydrophobic polymers to hydrophilic polymer-
        grafted carbon black and
        dispersibility)
IT
     Polymerization
        (graft; postgrafting of hydrophobic polymers to
        hydrophilic polymer-grafted carbon
        black and dispersibility)
TI
     Dispersion (of materials)
     Hydrophilicity
     Hydrophobicity
     Interface
        (postgrafting of hydrophobic polymers to hydrophilic polymer-
        grafted carbon black and
        dispersibility)
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IT 9002-98-6DP, reaction products with carbon black and glycidyl methacrylate-Me methacrylate copolymer 26141-88-8DP, Glycidyl methacrylatemethyl methacrylate copolymer, reaction products with carbon black and poly(ethyleneimine)

(postgrafting of hydrophobic polymers to hydrophilic polymer-grafted carbon black and

IT 538-75-0, N,N'-Dicyclohexylcarbodiimide

(postgrafting of hydrophobic polymers to hydrophilic polymergrafted carbon black and
dispersibility)

L30 ANSWER 11 OF 24 HCA COPYRIGHT 2006 ACS on STN 134:117224 Thermally-curable water-thinned coating command the control of th

dispersibility)

- 134:117224 Thermally-curable water-thinned coating compositions and multilayered coating films therefrom useful for metal and plastic protection. Masuda, Kazuaki; Osugi, Koji; Kuwashima, Teruaki; Harakawa, Takeshi (Nippon Paint Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 2001011151 A2 20010116, 16 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 2000-49321 20000225. PRIORITY: JP 1999-124008 19990430.
- AΒ The compns. having good water resistance and storage stability comprise a carboxy-contq. water-based polymer (A) and a hydrophilically modified polycarbodiimide (B) which has alternating units of carbodimides and alternating units of polyols linking to the previous units by urethane bondings and is terminated with hydrophilic units on 2 ends by urethane bondings. Thus, heating 700 parts 4,4-dicyclohexylmethane diisocyanate in the presence of 14 parts 3-methyl-1-phenyl-2-phospholene-1-oxide at 180° for 16 h, mixing 226.8 parts the resulting polycarbodiimide with 200 parts polypropylene glycol of Mn 2000 and heating at 90° for 3 h in the presence of 0.16 parts dibutyltin dilaurate gave an isocyanateterminated copolymer which was modified with polyoxyethylene mono(2ethylhexyl) ether to give a B-type copolymer. Mixing 80 parts a copolymer of Et acrylate 250, 2-hydroxyethyl methacrylate 150, 2hydroxyethyl acrylate 223, methacrylic acid 77 and styrene 300 parts with 20 parts B and 10 parts pigment paste gave a compn. which could be thinned with water.
- IT 24980-41-4DP, Polycaprolactone, diols, block copolymers with
 polycarbodiimide, modified with hydrophilic agents
 25248-42-4DP, Polycaprolactone, diols, block copolymers with
 polycarbodiimide, modified with hydrophilic agents

(thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

RN 24980-41-4 HCA

CN 2-Oxepanone, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 502-44-3 CMF C6 H10 O2

RN 25248-42-4 HCA
CN Poly[oxy(1-oxo-1,6-hexanediyl)] (9CI) (CA INDEX NAME)

IC ICM C08G018-79

ICS C08G018-83; C09D175-12

CC 42-10 (Coatings, Inks, and Related Products) Section cross-reference(s): 38, 55

acrylate polymer polycarbodiimide

polyalkylene glycol ether hydrophilic modifier

polycarbodiimide coating; water thinned hydrophilic modified

polycarbodiimide multiblock copolymer coating; thermosetting

hydrophilic modified polycarbodiimide multiblock copolymer

coating; dicyclohexylmethane diisocyanate polycarbodiimide

polyoxyethylene multiblock copolymer coating; polypropylene glycol

carbodiimides multiblock copolymer coating; carboxy contg

polymer polycarbodiimide alternating copolymer coating;

alternating copolymer coating; multilayered coating polycarbodiimide alternating copolymer

IT Phenoxy resins

(acrylic, coating vehicle; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

IT Polyesters, uses

Polyoxyalkylenes, uses

(polycarbodiimide-, block, hydrophilic component; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

IT Polyoxyalkylenes, uses

(polyester-, block, coating vehicle; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

IT Polycarbodiimides

(polyester-, block, hydrophilic component; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

IT Polyesters, uses

(polyoxyalkylene-, block, coating vehicle; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

IT Polycarbodiimides

(polyoxyalkylene-, block, hydrophilic component; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

TT 707-61-9, 3-Methyl-1-phenyl-2-phospholene-1-oxide (carbodimidization catalyst; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

77-99-6DP, Trimethylolpropane, alkyd resins IT 79-41-4DP, Methacrylic acid, polymers with acrylic and vinyl monomers and polyether -polysiloxanes 100-42-5DP, Styrene, polymers with acrylic and vinyl monomers and **polyether**-polysiloxanes 121-91-5DP, 124-04-9DP, Adipic acid, alkyd Isophthalic acid, alkyd resins resins 126-30-7DP, Neopentyl glycol, alkyd resins 140-88-5DP, Ethyl acrylate, polymers with acrylic and vinyl monomers and polyether-polysiloxanes 502-44-3DP, ε-Caprolactone, alkyd resins 552-30-7DP, Trimellitic 818-61-1DP, 2-Hydroxyethyl anhydride, alkyd resins acrylate, polymers with acrylic and vinyl monomers and polyether-polysiloxanes 868-77-9DP, 2-Hydroxyethyl methacrylate, polymers with acrylic and vinyl monomers and **polyether**-polysiloxanes 26915-97-9P 135991-20-7DP, Epol, alkyd resins 321181-75-3P, Ethyl acrylate-2-hydroxyethyl acrylate-2-hydroxyethyl methacrylatemethacrylic acid-styrene copolymer 321181-76-4P, Acrylamide-ethyl acrylate-2-hydroxyethyl methacrylate-methacrylic acid-methyl methacrylate copolymer 321181-77-5P, Carbonic acid-1,6-hexanediol-dimethylolpropionic acid-hydrazine-isophorone

acrylate; methacrylic acid-styrene **graft** copolymer (coating vehicle; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

diisocyanate block copolymer 321181-78-6P, Epikote EP-1256; ethyl

IT 13463-67-7, Tipaque R-820, uses

(pigment; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal

and plastic protection)

IT 53880-05-0DP, Isophorone diisocyanate polymer, reaction product polycarbonate diols and polyoxyethylene monolauryl ether 62948-28-1DP, 4,4'-Dicyclohexylmethane diisocyanate homopolymer, block copolymers with polycaprolactone diols, ethers with hydrophilic agents

(polycarbodiimide-contg.; thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

80-04-6DP, Hydrogenated bisphenol A, alkyd resins 9002-92-0DP, Polyethylene glycol monolauryl ether, reaction product with polycarbonate-polycarbodiimides 24980-41-4DP, Polycaprolactone, diols, block copolymers with polycarbodiimide, modified with hydrophilic agents 25248-42-4DP, Polycaprolactone, diols, block copolymers with polycarbodiimide, modified with hydrophilic agents 111460-07-2DP, Sodium hydroxypropanesulfonate, reaction products with polycaprolactone-polycarbodiimides

(thermally-curable water-thinned coating compns. and multilayered coating films therefrom useful for metal and plastic protection)

- L30 ANSWER 12 OF 24 HCA COPYRIGHT 2006 ACS on STN
- 133:267608 Resin composites with controlled phase structure of resins in the exfoliated surfaces of the fillers. Hasegawa, Naoki; Usuki, Arimitsu (Kabushiki Kaisha Toyota Chuo Kenkyusho, Japan). Ger. Offen. DE 10014254 Al 20000928, 18 pp. (German). CODEN: GWXXBX. APPLICATION: DE 2000-10014254 20000323. PRIORITY: JP 1999-78187 19990323.
- AB The composites comprise particles with surface area <50 m2/g and ≥2 polymers or a copolymer ≥2 types of segments. The polymers or the copolymer form/forms a phase structure with a unit thickness 1-1000 Nm and are/is on the surface of the exfoliated particles. It is preferential that the particles have a needle-shaped, plate- and(or) sheet-type, spherical, or cylindrical form and the copolymer is a block copolymer or a **graft** copolymer with ≥2 types of segments. A typical composite is formed from 400 g Tuftec H1013 and 38.4 g octadecylamine- montmorillonite adducts.
- IT **7440-44-0**, Carbon, uses

(particulates; resin composites with controlled phase structure of resins in exfoliated surfaces of fillers)

- RN 7440-44-0 HCA
- CN Carbon (7CI, 8CI, 9CI) (CA INDEX NAME)

```
ΙT
     9011-14-7, PMMA
        (resin composites with controlled phase structure of resins in
        exfoliated surfaces of fillers)
RN
     9011-14-7 HCA
CN
     2-Propenoic acid, 2-methyl-, methyl ester, homopolymer (9CI)
     INDEX NAME)
          1
    CM
    CRN 80-62-6
    CMF C5 H8 O2
IC
     ICM C08J003-20
     ICS C08J005-10; C09D005-00; B32B005-04; C08L101-12
     37-6 (Plastics Manufacture and Processing)
CC
ST
    nanocomposite graft copolymer controlled phase;
    octadecylamine montmorillonite adduct nanocomposite controlled
    polymer phase; polymer blend controlled phase nanocomposite
IT
    Polyesters, uses
        (arom.; resin composites with controlled phase structure of
        resins in exfoliated surfaces of fillers)
    Urethane rubber, uses
IT
        (polyester-, block, Elastollan C 95A; resin composites
        with controlled phase structure of resins in exfoliated surfaces
        of fillers)
ΙT
     Polyimides, uses
     Polyimides, uses
        (polyether-; resin composites with controlled phase
        structure of resins in exfoliated surfaces of fillers)
ΙT
     Polyamides, uses
     Polyamides, uses
      Polyethers, uses
      Polyethers, uses
        (polyimide-; resin composites with controlled phase structure of
        resins in exfoliated surfaces of fillers)
IT
    Acrylic polymers, uses
    Butyl rubber, uses
    EPDM rubber
     Fluoropolymers, uses
     Polyamides, uses
```

Polybenzimidazoles

Polycarbodiimides

Polycarbonates, uses

Polyesters, uses

Polyethers, uses

Polyimides, uses

Polyketones

Polyoxymethylenes, uses

Polysilanes

Polysiloxanes, uses

Polysulfones, uses

Polythiophenylenes

Polyureas

Polyurethanes, uses

Styrene-butadiene rubber, uses

(resin composites with controlled phase structure of resins in exfoliated surfaces of fillers)

IT **7440-44-0**, Carbon, uses

(particulates; resin composites with controlled phase structure of resins in exfoliated surfaces of fillers)

9002-84-0, PTFE 9002-88-4, Polyethylene 9003-17-2, Polybutadiene 9003-17-2D, Polybutadiene, hydrogenated 9003-29-6, Polybutene 9003-31-0, Polyisoprene 9003-31-0D, Polyisoprene, hydrogenated 9003-55-8D, Butadiene-styrene copolymer, hydrogenated 9010-79-1, Ethylene-propylene copolymer 9011-14-7, PMMA 9019-29-8, Butene-ethylene copolymer 9078-70-0, Polypentene 25212-74-2, Polythiophenylene

(resin composites with controlled phase structure of resins in exfoliated surfaces of fillers)

- L30 ANSWER 13 OF 24 HCA COPYRIGHT 2006 ACS on STN
- 132:294891 Resin compositions for use in finishing of cellulosic fabrics and method for their use. Mizushima, Makoto; Kaneida, Kenta (Nippon Shokubai Kagaku Kogyo Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 2000119968 A2 20000425, 11 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1998-290393 19981013.
- The resin compns. useful as binders for fabric printing inks, impregnation, coating, etc., comprise, as main compns., (A) polymers bearing COOH groups, and (B) polyfunctional oxazoline compds. or/and polyfunctional carbodiimide compds. as crosslinking agents, and (C) compds. bearing groups (X) which are reactive to OH groups and can form COOH groups by the reaction with OH groups, or/and compds. bearing the X groups and COOH groups for improving the adhesion of binders to fabric surfaces and retaining good printing color washfastness. Thus, prepg. a reducer from Bissurf 1400 (thickening emulsifier) 3, water 34 and mineral terpene 63 parts, mixing 75 parts the reducer with Ryudye W Blue KW (water-based pigment) 20, Nikasol A

- 01 (carboxy group-contg. acrylic polymer; 45.2% aq. soln.) 20, a Me methacrylate-2-isopropenyl-2-oxazoline-Light Acrylate 130A (methoxypolyethylene glycol acrylate) copolymer (40.4% aq. soln.) 2 and SMA 1000P (maleic anhydride-styrene copolymer) 2 parts gave an ink which was printed on a cotton fabric to give prints with good friction resistance and washfastness.
- IC ICM D06M015-263 ICS C08K005-29; C08K005-353; C08L035-00; C08L039-04; C08L079-00; C08L101-08; D06P001-46; D06P003-60
- CC 38-3 (Plastics Fabrication and Uses) Section cross-reference(s): 40
- ST textile printing ink binder acrylic resin oxazoline crosslinker; washfastness cotton fabric printing ink binder; wear resistance cotton fabric printing ink binder; cellulosic fabric finishing binder acrylic resin crosslinker oxazoline compd
- IT 25586-20-3P, Acrylic acid-butyl acrylate-styrene copolymer 25686-45-7P, Acrylic acid-acrylonitrile-butyl acrylate copolymer 264906-85-6P, Nikasol A 01

(base resins; resin compns. for use in finishing of cellulosic fabrics and method for use)

- IT 264266-13-9P, 2-Isopropenyl-2-oxazoline-Light Acrylate 130A-methacrylic acid **graft** copolymer 264266-14-0P, Ethylene oxide-2-isopropenyl-2-oxazoline-methacrylic acid **graft** copolymer 264266-15-1P, Acrylic acid-2-isopropenyl-2-oxazoline-styrene copolymer
 - (crosslinker; resin compns. for use in finishing of cellulosic fabrics and method for use)
- L30 ANSWER 14 OF 24 HCA COPYRIGHT 2006 ACS on STN
- 132:124272 Aqueous soil-repellent silicone polymer materials and their coating films. Tamura, Tsuruki; Tadaoka, Eisuke; Kawata, Hiroyuki (Hitachi Chemical Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 2000034422 A2 20000202, 6 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1998-204521 19980721.
- The coating films are obtained by reaction of polyisocyanates with the materials obtained from thermosetting polymer-type unsatd. esters, silicones RSiMe2[(OSiMe2)n(CH2)k(O)1(CH2)m]Z (I; R = Me, Ph; Z = OH, epoxy, CO2H, methacryloxy, acryloxy, dimethylolalkyl; n = 6-300; k = 1-10; l = 0-1; m = 0-6), and dicyclohexylcarbodiimide (II). Thus, reaction of 50 g vinyl ester (PS 6150; no.-av. mol. wt. 1650, OH value 131 mgKOH/g) with 58 g I (R = Me, Z = CO2H, n .simeq. 10, k = 10, l = m = 0; Silaplane FM 0611) in the presence of 14.5 g II in CHCl3 gave a graft copolymer, 10 parts of which was cured with 1 part HMDI (Coronate HX) on a stainless steel sheet to give a 10 μ m-thick film showing good antisoiling effect against soy sauce and water-repellent properties.

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IC
     ICM C09D005-00
     ICS C09D175-04; C09D183-10
CC
     42-10 (Coatings, Inks, and Related Products)
ST
     unsatd polyester silicone graft antisoiling
     coating; polyisocyanate crosslinked polyester siloxane
     antisoiling coating; water repellent coating polyester
     siloxane graft; dicyclohexylcarbodiimide
     graft polyester siloxane antisoiling coating
IT
     Coating materials
        (antisoiling; ag. soil-repellent coatings contg.
        polyisocyanate-crosslinked unsatd. polyester-silicone
        graft polymers)
IT
     Polymerization catalysts
        (graft, dicyclohexylcarbodiimide; aq.
        soil-repellent coatings contq. polyisocyanate-crosslinked unsatd.
        polyester-silicone graft polymers)
IT
     Polysiloxanes, uses
     Polysiloxanes, uses
        (polyester-, graft, polyisocyanate-
        crosslinked; aq. soil-repellent coatings contq.
        polyisocyanate-crosslinked unsatd. polyester-silicone
        graft polymers)
IT
     Polyesters, uses
       Polyesters, uses
        (polysiloxane-, graft, polyisocyanate-crosslinked; aq.
        soil-repellent coatings contq. polyisocyanate-crosslinked unsatd.
        polyester-silicone graft polymers)
IT
     Coating materials
        (water-resistant; aq. soil-repellent coatings contq.
        polyisocyanate-crosslinked unsatd. polyester-silicone
        graft polymers)
TΤ
     538-75-0, Dicyclohexylcarbodiimide
        (aq. soil-repellent coatings contq. polyisocyanate-crosslinked
        unsatd. polyester-silicone graft polymers)
ΙT
     256379-01-8P
                    256379-02-9P
                                   256382-03-3P 256382-04-4P
     256382-05-5P
                    256382-06-6P
        (aq. soil-repellent coatings contq. polyisocyanate-crosslinked
        unsatd. polyester-silicone graft polymers)
    ANSWER 15 OF 24 HCA COPYRIGHT 2006 ACS on STN
L30
           In vivo assessment of a novel dacron surface with covalently
     bound recombinant hirudin. Wyers, Mark C.; Phaneuf, Matthew D.;
     Rzucidlo, Eva M.; Contreras, Mauricio A.; LoGerfo, Frank W.; Quist,
     William C. (Division of Vascular Surgery, Beth Israel Deaconess
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Medical Center, Harvard Medical School, Children's Hospital, Boston, MA, 02215, USA). Cardiovascular Pathology, 8(3), 153-159 (English)

1999. CODEN: CATHE8. ISSN: 1054-8807. Publisher: Elsevier

Science Inc..

AΒ Prosthetic arterial graft surfaces are relatively thrombogenic and fail to heal with a cellular neointima. The goal of this study was to characterize the in vivo antithrombin properties of a novel Dacron surface with covalently linked recombinant hirudin (rHir) implanted in a canine thoracic aorta with high flow and shear rates. bound to a knitted Dacron patch using crosslinker-modified bovine serum albumin (BSA) as a basecoat protein. BSA was first reacted with the hetero-bifunctional crosslinker, sulfo-SMCC. This BSA-SMCC complex was then bound to the carboxylic acid groups of hydrolyzed Dacron patches using the carbodimide crosslinker, 1-ethyl-3-(3dimethylaminopropyl) carbodiimide hydrochloride. Iodinated, Traut'smodified rHir (125I-rHir-SH) was then reacted with the Dacron-BSA-SMCC surface, thereby covalently binding 125I-rHir. Graft segments were washed and sonicated to remove any nonspecifically bound 125I-Dacron-BSA-SMCC-S-125I-rHir patches (n = 5) and control Dacron-BSA patches (n = 5) were implanted in series in the thoracic aortas of canines. These patches were exposed to nonheparinized, arterial blood flow for 2 h. Patches were explanted and assessed for 125I-rHir loss. Antithrombin activity of explanted 1-cm2 patch segments was evaluated using a chromogenic assay with 1, 5, 10, 15 units of added thrombin. Light microscopy was performed to qual. examine the pseudointima. Two animals were excluded from the study owing to excessive bleeding through the knitted 125I-rHir patch. Comparison of preoperative and postoperative 125I-rHir y counts revealed an overall decrease of 20 ± 5.4% over the period studied. Explanted 125I-rHir patch segments were able to inhibit 1, 5, and 7 NIHU of thrombin, demonstrating retained antithrombin activity. Gross and microscopic examn. of the control and test Dacron surfaces showed marked differences. Dacron surfaces with covalently bound 125I-rHir had no gross thrombus and a thin pseudointima of platelets and plasma proteins. In contrast, the control patches had a thick pseudointima composed of fibrin rich thrombus. RHir, covalently bound to Dacron patches, maintains its biol. activity as well as prevents thrombus formation on the graft surface. This novel antithrombin coating, by modifying the blood/graft interface, may improve both short- and long-term patency in small-diam. prosthetic arterial grafts and has applications with respect to other implantable or indwelling biomaterials.

CC 63-7 (Pharmaceuticals)

Section cross-reference(s): 1

IT **Polyester** fibers, biological studies (conjugates with hirudin; in vivo assessment of novel dacron surface with covalently bound recombinant hirudin)

L30 ANSWER 16 OF 24 HCA COPYRIGHT 2006 ACS on STN 130:325734 A novel gas sensor from crystalline polymer-grafted

carbon black. Responsibility of electric resistance of composite from crystalline polymer-grafted carbon black against solvent vapor. Tsubokawa, Norio; Shirai, Yukio; Okazaki, Masaki; Maruyama, Kiyotaka (Department Material Science Technology, Faculty Engineering, Niigata University, Niigata, 950, Japan). Polymer Bulletin (Berlin), 42(4), 425-431 (English) 1999. CODEN: POBUDR. ISSN: 0170-0839. Publisher: Springer-Verlag. Cryst. polymers, such as $poly(\epsilon$ -caprolactone) (PCL), poly(ethylene)adipate) (PEA), and polyethylene (PE), were successfully grafted onto carbon black surface by direct condensation of terminal groups of these polymers with carboxyl groups on the surface using N, N'dicyclohexylcarbodiimide as a condensing agent. The elec. resistance of a composite prepd. from these cryst. polymer- grafted carbon black drastically increased to 104-105 times of initial resistance in good solvent vapor of grafted polymer and returned to initial resistance when it was transferred into dry air. However, the change of elec. resistance of the composite hardly obsd. in poor solvent vapor. These results suggest that these composite can be applied as a novel gas sensor. 24980-41-4DP, Poly(ϵ -caprolactone), graft products with surface-modified carbon black 25248-42-4DP, Poly[oxy(1-oxo-1,6-hexanediyl)], graft products with surface-modified carbon black (prepn. of cryst. polymer-grafted carbon blacks and elec. resistance of their composites usable as gas sensors) 24980-41-4 HCA 2-Oxepanone, homopolymer (9CI) (CA INDEX NAME) CM 1 502-44-3 CRN CMF C6 H10 O2

AΒ

ΙT

RN

CN

RN 25248-42-4 HCA CN Poly[oxy(1-oxo-1,6-hexanediyl)] (9CI) (CA INDEX NAME)

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[ ----- O- C- (CH2)5----- ]<sub>n</sub>
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CC
     37-5 (Plastics Manufacture and Processing)
     Section cross-reference(s): 76
ST
     elec resistance polymer grafted carbon
    black composite; gas sensor polymer grafted
     carbon black composite; polycaprolactone
    grafted carbon black gas
     sensor; polyethylene adipate grafted carbon
    black gas sensor; polyethylene grafted
    carbon black gas sensor
TI
    Natural gas, analysis
        (liquefied, vapor; gas sensor properties of cryst. polymer
        composites contq. grafted carbon
       black)
ΙT
    Carbon black, preparation
        (polymer-grafted; prepn. of cryst. polymer-
        grafted carbon blacks and elec.
        resistance of their composites usable as gas sensors)
ΙT
     Electric resistance
     Gas sensors
        (prepn. of cryst. polymer-grafted carbon
       blacks and elec. resistance of their composites usable as
        gas sensors)
ΙT
    Kerosene
    Ligroine
        (vapor; gas sensor properties of cryst. polymer composites contg.
        grafted carbon black)
     9002-88-4P, Polyethylene 24937-05-1P, Poly(ethylene adipate), SRU
IT
     24938-37-2P, Poly(ethylene adipate) 24980-41-4DP,
     Poly(ε-caprolactone), graft products with
     surface-modified carbon black
     25248-42-4DP, Poly[oxy(1-oxo-1,6-hexanediyl)], graft
     products with surface-modified carbon black
        (prepn. of cryst. polymer-grafted carbon
       blacks and elec. resistance of their composites usable as
        gas sensors)
IT
     67-56-1, Methanol, analysis 74-98-6, Propane, analysis
     Butane, analysis 108-88-3, Toluene, analysis
                                                     109-99-9, THF,
                                             141-78-6, Ethyl acetate,
     analysis 110-54-3, Hexane, analysis
     analysis 7732-18-5, Water, analysis
        (vapor; gas sensor properties of cryst. polymer composites contg.
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grafted carbon black)

L30 ANSWER 17 OF 24 HCA COPYRIGHT 2006 ACS on STN
127:359260 Nontoxic and safe polycarbodiimides, their
preparation and use as crosslinking agents, curable at low
temperatures, in various resin compositions. Nakamura, Michiei;
Simanaka, Hiroyuki; Sugawara, Eiichi; Wakebe, Yoshitaka; Okura, Ken;
Kawamura, Tatsuo; Takahashi, Masayuki; Takezawa, Nobuo
(Dainichiseika Color and Chemicals Mfg. Co., Ltd., Japan; Ukima
Colour & Chemicals Mfg. Co., Ltd.). Eur. Pat. Appl. EP 805172 A2
19971105, 11 pp. DESIGNATED STATES: R: CH, DE, ES, FR, GB,
IT, LI. (English). CODEN: EPXXDW. APPLICATION: EP 1997-106927

PRIORITY: JP 1996-130624 19960430.

AΒ A polyfunctional polycarbodiimide compd., which comprises at least four mol. chains contq. a carbodiimido N:C:N group bonded independently to a backbone, can be produced by reacting (a) an isocyanate compd. having at least one carbodiimido group and at least one isocyanate group with (b) a polyol, polyamine and/or amino alc. having at least four hydroxyl, primary amino and/or secondary amino groups in a mol. Thus, 631.4 parts (by wt.) 30% soln. of polyhexamethylenecarbodiimide diisocyanate obtained by condensing 4 mols. of hexamethylene diisocyanate using a carbodiimidation catalyst was mixed with 1.3 parts 5% soln. of dibutyltin dilaurate and 468.5 parts 50% soln. of polyethylene glycol monomethyl ether, followed by an addn. of 37.6 parts 50% soln. of decaglyceryl monolaurate to give a polycarbodiimide compd. (crosslinking agent) having 11 carbodiimido-contg. side chains and 30 carbodiimido groups in total in a mol. A resin-based printing paste comprising 4:60:36 acrylic acid-Et acrylate-styrene copolymer latex 20, 20% solids crosslinking agent prepd. above 5, water 10, 20% solids polyoxyethylene alkylphenyl ether 5, mineral terpin 55, and aq. copper phthalocyanine blue pigment 5 parts was applied on a knitted cotton fabric by a screen printing machine and the printed fabric was excellent in various fastnesses, soft, and a vivid blue color.

IT 25248-42-4D, Poly[oxy(1-oxo-1,6-hexanediyl)], diol derivs., polymers with diols and TDI

(prepn. of **polycarbodiimide** crosslinking agents and their use in low-temp.-curable resin compns. for treatment of articles)

RN 25248-42-4 HCA

19970425.

CN Poly[oxy(1-oxo-1,6-hexanediyl)] (9CI) (CA INDEX NAME)

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IC
     ICM C08G018-79
     ICS C08G018-10; C08G018-28; C08G018-08; C07C267-00; C08G085-00;
          C08L101-00
ICI
    C08L101-00, C08L079-00
CC
     35-5 (Chemistry of Synthetic High Polymers)
     Section cross-reference(s): 39, 40, 42, 43, 55
ST
     nontoxic polycarbodiimide crosslinking agent prepn; safety
     resin compn polycarbodiimide crosslinking agent;
     hexamethylene diisocyanate decaglyceryl monolaurate copolymer prepn;
     polyethylene glycol monomethyl ether termination
     polycarbodiimide; acrylic acid ethyl acrylate styrene
     copolymer; printing cotton fabric knit
ΙT
     Automobiles
        (bumpers, polypropylene, moldings; prepn. of
        polycarbodiimide crosslinking agents and their use in
        low-temp.-curable resin compns. for treatment of articles)
IT
     Acrylic polymers, uses
       Polyesters, uses
     Polyurethanes, uses
        (carboxy-contg.; prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
        compns. for treatment of articles)
IT
     Textiles
        (cotton, knitted, printing pastes for; prepn. of
        polycarbodiimide crosslinking agents and their use in
        low-temp.-curable resin compns. for treatment of articles)
IT
     Polyester fibers, uses
        (fabrics, tufted or woven, coatings for; prepn. of
        polycarbodiimide crosslinking agents and their use in
        low-temp.-curable resin compns. for treatment of articles)
IT
     Polyamides, uses
       Polyesters, uses
        (films, inks for; prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
        compns. for treatment of articles)
IT
     Inks
        (gravure, water-thinned; prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
        compns. for treatment of articles)
ΙΤ
     Inks
        (gravure; prepn. of polycarbodiimide crosslinking
        agents and their use in low-temp.-curable resin compns. for
        treatment of articles)
ΙT
    Acrylic polymers, uses
        (hydroxy-contg.; prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
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compns. for treatment of articles)
ΙT
     Plastic films
        (inks for gravure printing of; prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
        compns. for treatment of articles)
ΙT
     Coating materials
        (moisture-permeable; prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
        compns. for treatment of articles)
IT
     Textile printing
     Textile printing
        (pastes; prepn. of polycarbodiimide crosslinking agents
        and their use in low-temp.-curable resin compns. for treatment of
        articles)
ΙT
     Floors
        (poly(vinyl chloride); prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
        compns. for treatment of articles)
IT
     Polyoxyalkylenes, preparation
     Polyoxyalkylenes, preparation
     Polyoxyalkylenes, preparation
        (polycarbodiimide-polyurethane-, crosslinking agent;
        prepn. of polycarbodiimide crosslinking agents and
        their use in low-temp.-curable resin compns. for treatment of
        articles)
     Polyurethanes, preparation
ΙT
     Polyurethanes, preparation
     Polyurethanes, preparation
        (polyoxyalkylene-polycarbodiimide-, crosslinking agent;
        prepn. of polycarbodiimide crosslinking agents and
        their use in low-temp.-curable resin compns. for treatment of
        articles)
IT
     Polycarbodiimides
       Polycarbodiimides
       Polycarbodiimides
        (polyoxyalkylene-polyurethane-, crosslinking agent; prepn. of
        polycarbodiimide crosslinking agents and their use in
        low-temp.-curable resin compns. for treatment of articles)
ΙT
     Polyurethanes, uses
     Polyurethanes, uses
        (polyurea-; prepn. of polycarbodiimide crosslinking
        agents and their use in low-temp.-curable resin compns. for
        treatment of articles)
ΙT
     Polyureas
     Polyureas
        (polyurethane-; prepn. of polycarbodiimide crosslinking
        agents and their use in low-temp.-curable resin compns. for
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treatment of articles)
ΙT
     Adhesives
     Coating materials
     Crosslinking agents
       Paints
     Waterproofing agents
        (prepn. of polycarbodiimide crosslinking agents and
        their use in low-temp.-curable resin compns. for treatment of
        articles)
IT
     Laminated plastic films
        (prepn. of polycarbodiimide crosslinking agents and
        their use in low-temp.-curable resin compns. for treatment of
        articles)
ΙT
     Polyurethanes, preparation
        (prepn. of polycarbodiimide crosslinking agents and
        their use in low-temp.-curable resin compns. for treatment of
        articles)
IT
    Metals, uses
     Nonwoven fabrics
     Paper
     Plastics, uses
     Textiles
     Threads
     Wood
     Yarns
        (prepn. of polycarbodiimide crosslinking agents and
        their use in low-temp.-curable resin compns. for treatment of
        articles)
ΙT
     Acrylic rubber
        (prepn. of polycarbodiimide crosslinking agents and
        their use in low-temp.-curable resin compns. for treatment of
        articles)
IT
     Inks
        (printing; prepn. of polycarbodiimide crosslinking
        agents and their use in low-temp.-curable resin compns. for
        treatment of articles)
IT
     Pastes
     Pastes
        (textile printing; prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
        compns. for treatment of articles)
ΙT
     Coating materials
        (water-resistant; prepn. of polycarbodiimide
        crosslinking agents and their use in low-temp.-curable resin
        compns. for treatment of articles)
IT
    Adhesives
        (water-thinned; prepn. of polycarbodiimide crosslinking
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agents and their use in low-temp.-curable resin compns. for treatment of articles)

71-36-3DP, 1-Butanol, reaction products with polyoxyalkylene-ΙT polycarbodiimide-polyurethanes, preparation 143-28-2DP, reaction products with polyoxyalkylene-polycarbodiimide -polyurethanes 9003-13-8DP, Polypropylene glycol monobutyl ether, reaction products with polyoxyalkylene-polycarbodiimide -polyurethanes 9004-74-4DP, Polyethylene glycol monomethyl ether, reaction products with polyoxyalkylene-polycarbodiimide 9038-95-3DP, Polyoxyethylene-polyoxypropylene -polyurethanes monobutyl ether, reaction products with polyoxyalkylenepolycarbodiimide-polyurethanes 198636-00-9DP, Dipentaerythritol monolaurate-hexamethylene diisocyanate copolymer, reaction products with polypropylene glycol monobutyl ether 198646-64-9DP, Decaglycerol monolaurate-hexamethylene diisocyanate copolymer, reaction products with polyethylene glycol monomethyl 198646-65-0DP, Hexamethylene diisocyanate-polyoxyethylene ether sorbitol monolaurate copolymer, reaction products with polyoxyethylene-polyoxypropylene monobutyl ether or polyethylene glycol monomethyl ether 198646-66-1DP, Polyoxyethylene sorbitol monolaurate-tolylene diisocyanate copolymer, reaction products with polyethylene glycol monomethyl ether, oleyl alc., n-butanol, or n-butanol and sulfanyltriethylamide

(crosslinking agent; prepn. of **polycarbodiimide** crosslinking agents and their use in low-temp.-curable resin compns. for treatment of articles)

IT 9002-86-2, Poly(vinyl chloride)

(film or flooring material, inks or coatings for; prepn. of **polycarbodiimide** crosslinking agents and their use in low-temp.-curable resin compns. for treatment of articles)

IT 9003-07-0, Polypropylene

(film or molding, adhesives and inks and coatings for; prepn. of polycarbodiimide crosslinking agents and their use in

low-temp.-curable resin compns. for treatment of articles)

IT 9002-88-4, Polyethylene

(film, inks for; prepn. of **polycarbodiimide** crosslinking agents and their use in low-temp.-curable resin compns. for treatment of articles)

107896-64-0P, Diphenylmethane diisocyanate-ethylene oxide-propylene oxide block copolymer 198636-01-0P, Dimethylolpropionic acid-diphenylmethane diisocyanate-ethylene glycol-tetramethylene glycol copolymer

(prepn. of **polycarbodiimide** crosslinking agents and their use in low-temp.-curable resin compns. for treatment of articles)

IT 629-11-8D, 1,6-Hexanediol, polymers with diols and TDI 4767-03-7D, polymers with diols and TDI 25119-83-9, Acrylic acid-butyl

acrylate copolymer 25135-39-1, Acrylic acid-ethyl acrylate-methyl methacrylate copolymer 25215-62-7, Monobutyl maleate-styrene copolymer 25248-42-4D, Poly[oxy(1-oxo-1,6-hexanediyl)], diol derivs., polymers with diols and TDI 25585-77-7, Acrylic acid-ethyl acrylate-styrene copolymer 26471-62-5D, TDI, polymers with polycaprolactonediol and polyols 29035-81-2, Acrylic acid-vinyl acetate-vinyl chloride copolymer 198636-03-2, Diethylenetriamine-dimethylolpropionic acid-isophorone diisocyanate-tetramethylene glycol copolymer triethylamine salt 198636-04-3, Ethyl methacrylate-2-hydroxyethyl methacrylate-methacrylic acid-methyl methacrylate-octyl methacrylate copolymer

(prepn. of **polycarbodiimide** crosslinking agents and their use in low-temp.-curable resin compns. for treatment of articles)

- L30 ANSWER 18 OF 24 HCA COPYRIGHT 2006 ACS on STN
- 127:39703 Covalent linkage of recombinant hirudin to poly(ethylene terephthalate) (Dacron): creation of a novel antithrombin surface. Phaneuf, Matthew D.; Berceli, Scott A.; Bide, Martin J.; Quist, William C.; LoGerfo, Frank W. (Vascular Surgery Research, Deaconess Hospital/Harvard Medical School, Boston, MA, 02215, USA). Biomaterials, 18(10), 755-765 (English) 1997. CODEN: BIMADU. ISSN: 0142-9612. Publisher: Elsevier.
- AΒ Thrombus formation and intimal hyperplasia on the surface of implantable biomaterials such as poly(ethylene terephthalate)(Dacron) vascular grafts are major concerns when utilizing these materials in the clin. setting. Thrombin, a pivotal enzyme in the blood coagulation cascade primarily responsible for thrombus formation and smooth muscle cell activation, has been the target of numerous strategies to prevent this phenomenon from occurring. The purpose of this study was to covalently immobilize the potent, specific antithrombin agent recombinant hirudin (rHir) to a modified Dacron surface and characterize the in vitro efficacy of thrombin inhibition by this novel biomaterial surface. Bovine serum albumin (BSA), which was selected as the 'basecoat' protein was reacted with various molar ratios of the crosslinker sulfosuccinimidyl 4-(N-maleimidomethyl) cyclohexane-1-carboxylate (sulfo-SMCC; 1:5-1:50). These BSA-SMCC complexes were then covalently linked to sodium hydroxide-hydrolyzed Dacron (HD) segments via the cross-linker 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDC). Covalent linkage of these complexes to HD (HD-BSA-SMCC) was not affected by any of the sulfo-SMCC cross-linker ratios assayed. RHir, which as initially reacted with 2-iminothiolane hydrochloride (Traut's reagent) in order to create sulphydryl groups, was then covalently bound to these HD-BSA-SMCC surfaces (HD-BSA-SMCC-S-rHir). The 1:50 (BSA:sulfo-SMCC) HD-BSA-SMCC-S-rHir segments bound 22-fold more rHir (111 bg oer ng

/Dacron) compared to control segments and also possessed the greatest thrombin inhibition of the segments evaluated using a **chromogenic** substrate assay for thrombin. Further characterization of the HD-BSA-SMCC-S-rHir segments demonstrated that max. thrombin inhibition was 20.43 NIHU, 14.6-fold greater inhibition than control segments (1.4 NIHU). Thrombin inhibition results were confirmed by 125I-thrombin binding expts., which demonstrated that the 1:50 HD-BSA-SMCC-S-rHir segments was also significantly less than the control segments. Thus, these results demonstrate that rHir can be covalently bound to a clin. utilized biomaterial (Dacron) while still maintaining its ability to bind and inhibit thrombin.

- CC 63-7 (Pharmaceuticals)
- ST hirudin conjugate polyethylene terephthalate antithrombogenic; vascular **graft** hirudin conjugate Dacron
- IT **Polyesters**, biological studies (reaction products with hirudin; covalent linkage of recombinant hirudin to poly(ethylene terephthalate) for a antithrombin surface)
- L30 ANSWER 19 OF 24 HCA COPYRIGHT 2006 ACS on STN
- 124:298767 Surface modification of polyetherurethaneureas and their antithrombogenicity. Kang, I.-K.; Kwon, O. H.; Byun, K. H.; Kim, Y. H. (Dep. Polymer Sci., Kyungpook Natl. Univ., Taegu, 702, S. Korea). Journal of Materials Science: Materials in Medicine, 7(3), 135-40 (English) 1996. CODEN: JSMMEL. ISSN: 0957-4530. Publisher: Chapman & Hall.
- AΒ Polyetherurethaneurea (PU) films were treated by oxygen plasma discharge followed by acrylic acid (AA) grafting. The carboxyl groups of the AA-grafted PU (PU-AA) surface were coupled with bovine serum albumin and heparin via water sol. carbodiimide. characterization of the modified PUs was carried out by attenuated total reflection Fourier transform IR (ATR-FTIR) spectroscopy and electron spectroscopy for chem. anal. (ESCA). The amt. of immobilized albumin (AL) and heparin (HE) on the PU surface was 1.8 and 1.5 μ g/cm2, resp., as detd. by the **dye** interaction method. Interactions between the surface-modified PUs and blood components such as plasma proteins and platelets were investigated to evaluate the blood compatibility of the samples. Plasma recalcification time (PRT) and activated partial thromboplastin time (APTT) of the albumin-immobilized PU (PU-AL) were almost the same as those of PU, while platelets were less adhered on the PU-Al than on PU. On the other hand, PRT and APTT of the PU-HE were significantly longer than those of the PU, PU-AA, and PU-AL. Moreover, adhesion of platelets was effectively suppressed on the PU-HE, leading to good in vitro blood compatibility.
- CC 63-7 (Pharmaceuticals)
 Section cross-reference(s): 38

- ST **polyether** urethane urea surface modification antithrombogenicity; antithrombogenic prosthetic polyetherurethaneurea surface modification

- 79-10-7DP, Acrylic acid, **grafts** with polyetherurethaneurea 9053-66-1DP, 4,4'-Diphenylmethane diisocyanate-ethylene diamine-polytetramethylene glycol copolymer, **grafts** with acrylic acid

(surface modification effect on antithrombogenicity of polyetherurethaneureas)

- L30 ANSWER 20 OF 24 HCA COPYRIGHT 2006 ACS on STN

 123:315561 Grafting reaction of surface carboxyl groups on
 carbon black with polymers having terminal
 hydroxyl or amino groups using N,N'-dicyclohexyl carbodiimide as a condensing agent. Tsubokawa, N.; Hosoya,
 M.; Kurumada, J. (Department of Material and Chemical Engineering,
 Faculty of Engineering, Niigata University, 8050, Ikarashi 2-nocho,
 Niigata, 950-21, Japan). Reactive & Functional Polymers, 27(1),
 75-81 (English) 1995. CODEN: RFPOF6. ISSN: 1381-5148.
 Publisher: Elsevier.
- AΒ Grafting of polymers onto carbon black [Neospectra II, Columbian Carbon Co.; channel black FW 200, Degussa A.G.; furnace black Philblack O, Phillips Petroleum Co. | surfaces was carried out by direct condensation of surface carboxyl groups with functional polymers. The reaction of surface carboxyl groups with functional polymers having hydroxyl or amino groups proceeded readily in the presence of N,N'-dicyclohexyl-carbodiimide (DCC) as condensing agent at 30° and the corresponding polymers were **grafted** onto **carbon black** surface with ester or amide bonds. The percentage of grafting of diol poly(propylene oxide) (PPG Mn = 2.0 + 103) and diamine-type poly(dimethylsiloxane) (SDA: Mn = 1.7 + 103) was 24.5 and 40.2%, resp. No grafting reaction onto carbon black surface, however, was obsd. in the absence of DCC. The percentage of grafting increased with increasing carboxyl group content of carbon black and increasing reaction temp. The percentage of grafting and the no. of grafted polymer chains decreased with increasing mol. wt. of functional polymers, because the reaction of surface carboxyl groups was inhibited by the already grafted polymer chains. The polymer-grafted carbon black gave a stable colloidal dispersion in a suitable solvent for the grafted polymer.

IT **25322-69-4DP**, Poly(propylene glycol), reaction products with carbon blacks

(grafting carbon black with

hydroxyl or amino terminated polymers using N,N'-dicyclohexyl-carbodiimide as a condensing agent)

RN 25322-69-4 HCA

ΙT

CN Poly[oxy(methyl-1,2-ethanediyl)], α -hydro- ω -hydroxy-(9CI) (CA INDEX NAME)

$$HO - (C3H6) - O - n$$

CC 37-6 (Plastics Manufacture and Processing)
Section cross-reference(s): 57

ST carbon black surface grafting polyglycol; polydiamine carbon black condensation imide

IT Siloxanes and Silicones, preparation

(diol- and diamine-terminated reaction products with carbon blacks; grafting
carbon black with hydroxyl or amino terminated polymers using N,N'-dicyclohexyl-carbodiimide as a

condensing agent)
Carbon black, preparation

(surface reaction products with diol- and diamine-terminated polymers; grafting carbon black with hydroxyl or amino terminated polymers using N,N'-dicyclohexyl-carbodiimide as a condensing agent)

IT 538-75-0, N,N'-Dicyclohexyl-carbodimide

(grafting carbon black with

hydroxyl or amino terminated polymers using N,N'-dicyclohexyl-carbodiimide as a condensing agent)

9016-00-6DP, Di-Me siloxane, SRU, reaction products with carbon blacks 25322-68-3DP, Poly(ethylene glycol), reaction products with carbon blacks

25322-69-4DP, Poly(propylene glycol), reaction products with carbon blacks 31900-57-9DP, Dimethylsilanediol

homopolymer, reaction products with carbon blacks

(grafting carbon black with

hydroxyl or amino terminated polymers using N,N'-dicyclohexyl-carbodimide as a condensing agent)

L30 ANSWER 21 OF 24 HCA COPYRIGHT 2006 ACS on STN

- 122:189197 Surface modification of carbon microbead by the **grafting** of polymers. Tsubokawa, Norio; Hayashi, Shinji (Faculty Engineering, Niigata University, Niigata, 950-21, Japan). Journal of Macromolecular Science, Pure and Applied Chemistry, A32(3), 525-35 (English) **1995**. CODEN: JSPCE6. ISSN: 1060-1325. Publisher: Dekker.
- AB The surface grafting of polymers onto carbon microbead, sphere of carbonized phenol resin, was achieved by two methods: (1) the radical graft polymn. initiated by azo groups introduced onto the surface and (2) the trapping of polymer radicals by a carbon microbead surface. The introduction of azo groups onto a carbon microbead surface was achieved by the reaction of surface carboxyl groups on the bead with 2,2'-azobis[2-(hydroxymethyl)- propionitrile] using N,N'dicyclohexylcarbodiimide as a condensing agent. The radical polymn. of Me methacrylate was initiated by surface azo groups introduced onto the surface, and PMMA was grafted onto the bead through the propagation of polymer chains from the surface: the percentage of grafting reached to 18% after 16 h. Furthermore, it was found that by the reaction of an azo polyamide (a polyamide having an azo bond in the main chain) and peroxide polymers (polymers having peroxide groups in the main chain and/or pendant group) with carbon microbead at 70-110°, polymer radicals formed by the thermal decompn. of these azo and peroxide polymers gave the corresponding polymer-grafted carbon microbead. The percentage of grafting decreased with increasing mol. wt. of the polymer radicals. The reactivity of surface functional groups and polycondensed arom. rings on carbon microbead are analogous with those of a carbon black surface.
- CC 37-6 (Plastics Manufacture and Processing)
- ST surface modification carbon microbead grafting; azo carbon microbead grafting polymer; methacrylate polymer grafting carbon microbead; polyamide azo grafting carbon microbead; peroxide polymer grafting carbon microbead

- IT 161776-43-8P 161776-44-9P 161776-45-0P (surface modification of carbon microbead by **grafting** of polymers)
- L30 ANSWER 22 OF 24 HCA COPYRIGHT 2006 ACS on STN

119:161523 Grafting of a functionalized sidechain liquid-crystal polymer on carbon fiber surfaces: novel coupling agents for fiber/polymer matrix composites. Le Bonheur, Vassoudevane; Stupp, Samuel I. (Dep. Mater. Sci. Eng., Univ. Illinois, Urbana, IL, 61801, USA). Chemistry of Materials, 5(9), 1287-92 (English) 1993. CODEN: CMATEX. ISSN: 0897-4756. AΒ Covalent grafting to functionalized carbon fibers of a specially designed liq.-cryst. monomer and its corresponding side-chain liq.cryst. polymer contg. pendant chem. functions on their mesogenic groups was studied. From a materials point of view these lig.-cryst. compds. could act as coupling agents as fiber/polymer matrix interfaces, offering a mechanism to control composite properties not only through bonding but also through their spontaneous mol. The grafting methodol. for both orientation in interfacial regions. monomer and polymer to fiber surfaces involved esterification through carbodiimide chem. in soln. Carboxylic acid groups found on functionalized carbon fiber surfaces were esterified to phenolic functions in the side chains of the exptl. polymer. Following grafting procedures the fibers were analyzed by SEM and by contact angle measurements. SEM micrographs of fibers grafted with polymer revealed the presence of strongly attached polymeric material on the graphitic surface after rigorous extn. with polymer solvent. Contact angle measurements and polar/dispersive free energy anal. indicated also a smaller polar component of the surface free energy of fibers possibly due to the hydrophobic polymer backbone grafted on the carbon surfaces. The esterification reaction grafted the polyphenolic liq.-crystal polymer on graphite fiber surfaces.

IT 7440-44-0

(carbon fibers, **grafting** on functionalized, with hydroxybiphenylbenzoate-contg. monomer and polymer liq. crystals, surface properties in relation to)

RN 7440-44-0 HCA

CN Carbon (7CI, 8CI, 9CI) (CA INDEX NAME)

c Ì

Folyloxy(methyl-1,2-ethanedlyl)], α -nydro- ω -nyd (9CI) (CA INDEX NAME)

$$HO - (C3H6) - O - H$$

- CC 37-6 (Plastics Manufacture and Processing) Section cross-reference(s): 75 STliq crystal grafting carbon fiber; surface property grafting carbon fiber; morphol grafting carbon fiber; contact angle grafting carbon fiber ΙT Carbon fibers, miscellaneous (grafting on functionalized, with hydroxybiphenylbenzoate-contg. monomer and polymer liq. crystals, surface properties in relation to) ΙT Liquid crystals, polymeric (hydroxybiphenylbenzoate deriv. polymers, functionalized carbon fiber grafting with, surface properties in relation to) ΙT Liquid crystals (hydroxybiphenylbenzoate derivs., functionalized carbon fiber grafting with, surface properties in relation to) ΙT Surface energy (of hydroxybiphenylbenzoate-contg. liq. crystal-grafted carbon fibers) ΙT Contact angle (of liqs., on hydroxybiphenylbenzoate-contg. liq. crystalgrafted carbon fibers) ΙT 7440-44-0 (carbon fibers, grafting on functionalized, with hydroxybiphenylbenzoate-contg. monomer and polymer liq. crystals, surface properties in relation to) IT 56-81-5, 1,2,3-Propanetriol, properties 75-12-7, Formamide, 544-76-3, n-Hexadecane 7732-18-5, Water, properties properties 25322-69-4, Polypropylene glycol (contact angle of, on hydroxybiphenylbenzoate-contg. liq. crystal-grafted carbon fibers) IT538-75-0, 1,3-Dicyclohexylcarbodiimide (grafting of hydroxybiphenylbenzoate-contq. liq. crystals on carbon fibers in presence of) ΙT 149828-91-1 149828-92-2 (grafting of lig.-cryst., on functionalized carbon
- L30 ANSWER 23 OF 24 HCA COPYRIGHT 2006 ACS on STN
 114:118095 Process for covalent surface modification of hydrophobic polymers and affinity membranes made therefrom. Azad, A. R. M.;
 Goffe, Randal A. (Sepracor, Inc., USA). PCT Int. Appl. WO 9004609

fibers, surface properties in relation to)

Al 19900503, 145 pp. DESIGNATED STATES: W: AU, BB, BG, BR, DK, FI, HU, JP, KR, LK, MC, MG, MW, NO, RO, SD, SU; RW: AT, BE, BF, BJ, CF, CG, CH, CM, DE, FR, GA, GB, IT, LU, ML, MR, NL, SE, SN, TD, TG. (English). CODEN: PIXXD2. APPLICATION: WO 1989-US4620 19891016. PRIORITY: US 1988-258406 19881017. The title process comprises e.g. (1) contacting a hydrophobic polymer with a soln. of a 1st nonsolubilizing solvent and a linker for sufficient time to form a covalent bond between the linker for sufficient time to form a covalent bond between the linker and a functionalizable side chain of the hydrophobic polymer; (2) contacting the reacted polymer of 1 with a soln. of a 2nd nonsolubilizing solvent and a mammol. for sufficient time to covalently bind the macromol. to the covalently bonded linker moiety. The above product may then be reacted with a reagent capable of producing active sites on the covalently bonded macromol., followed by reaction the produced active sites with a ligand. The process is conveniently carried out under heterogeneous conditions and proceeds with without a significant redn. in microporous membrane pore dimensions or hydraulic permeability of the original unmodified membrane. Also provided are a 4-component dope compn. and a spinnerette assembly useful for the manuf. of the polymers of the invention. Thus, poly(ether sulfone)/poly(ethylene oxide) hollow fiber membranes were prepd., treated with ethylene glycol diglycidyl ether, and then reacted with hydroxyethyl cellulose. fibers were activated with 2-fluoro-1-methylpyridinium ptoluenesulfonate and then reacted with antibodies to blood coagulation factor VIII. The resulting affinity membrane was used to purify factor VIII 115-fold from a factor VIII conc. Details of manuf. of the polymers of the invention are given, as are schematic diagrams of the spinnerette assembly. ICM C08F008-00 C08F008-06; C08F008-10; C08F008-14; C08F008-28; C08F008-32; C08F008-42; C08F008-44; C08F008-46; C08J003-24; C08J005-18; C08J005-20 9-3 (Biochemical Methods) Section cross-reference(s): 35 Polyoxyalkylenes, biological studies Acrylic polymers, uses and miscellaneous (in surface-modified hollow-fiber membrane manuf.) Polycarbonates, biological studies Polyesters, biological studies Polyimides, uses and miscellaneous Polymers, uses and miscellaneous Siloxanes and Silicones, biological studies Polysulfones, uses and miscellaneous Polythioarylenes

(surface modification of, for affinity membrane manuf.)

AΒ

IC

CC

IT

IT

ITDyes

Plasmid and Episome Surfactants Agglutinins and Lectins Antibodies Antigens Blood-coagulation factors Hormones Receptors

> (surface-modified polymer conjugate with, for affinity membrane manuf.)

IT Polyketones

L30

- Polysulfones, uses and miscellaneous (polyether-, surface modification of, for affinity membrane manuf.)
- ΙT Polyethers, biological studies (polyketone-, surface modification of, for affinity membrane
- IT Polyethers, uses and miscellaneous (polysulfone-, surface modification of, for affinity membrane
- 151-51-9, Carbodiimide 506-68-3, Cyanogen bromide IT 2224-15-9, Ethylene glycol diglycidyl ether 2425-79-8 58086-67-2, 2-Fluoro-1-methylpyridinium p-toluenesulfonate 67-63-0, 2-Propanol, uses and miscellaneous 75-05-8, Acetonitrile, 79-11-8, Chloroacetic acid, uses and uses and miscellaneous miscellaneous 106-89-8, uses and miscellaneous (in polymer surface modification for affinity membrane manuf.)
- ANSWER 24 OF 24 HCA COPYRIGHT 2006 ACS on STN 108:114377 Manufacture of fire- and chemically resistant elastomer compositions. Corish, Patrick Joseph; Ohbi, Daljit Singh (BICC PLC, Eur. Pat. Appl. EP 251792 A2 19880107, 10 pp. DESIGNATED STATES: R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU,
 - (English). CODEN: EPXXDW. APPLICATION: EP 1987-305854 NL, SE. 19870701. PRIORITY: GB 1986-16136 19860702.
- AΒ Halogen-free, fire-resistant, radiation-curable elastomeric compns., resistant to seawater, hydrocarbon oils, and hydraulic fluids, useful as elec. insulators for cables, comprise oil-resistant pigmented polyester-polyether and/or polyamide-polyether thermoplastic elastomers, nitrile rubber, optionally ≤1 part other diene or acrylic rubbers per 3 parts nitrile rubber, and Al203.3H2O. The nitrile rubber has a hydrocarbon main chain having ≤2 (epoxidized) double bonds per 5 C atoms, ≤1 CN side chains per 22 C atoms, and ≤1 hetero side chain per 4 C atoms (with ≥70% of the side chains being CN). The thermoplastic elastomer constitutes 10-70% of the total elastomer

content, and the Al2O3.3H2O content is enough to give an O index ≥ 25 but ≤2 times the total elastomer content. Thus, Hytrel 4056 (segmented polyester-polyether thermoplastic elastomer) 36, Chemigum NX 775 (carboxylated nitrile rubber) 60, Hydral 710B (Al203.3H20) 120, Hytrel 10 MS (master batch of polycarbodiimide in Hytrel) 4, SRF Black 10, 50% Silane A 172 4, Kemgard 911A (smoke suppressant) 4, 70% triallyl cyanurate 4, Dythal (Pb phthalate paste) 6, Monoplas 530 (plasticizer) 10, and Agerite Resin D 2 parts were mixed in 2 stages in a Banbury mixer, dumped, sheeted off, and cured by electron beam irradn. (dosage 175 kGy) to give specimens showing tensile strength 10.8 MPa, 100% modulus 9.6 MPa, elongation 250%, tear strength 5.6 MPa/mm, K factor insulation resistance 0.1 M Ω/km , and 0 index 30.5. When cured samples were exposed for 28 days to diesel fuel at 20°, hydraulic fluid OX-38 at 50°, and deionized H2O at 50°, they showed vol. swelling 8.4%, 6.3%, and 11%, resp., tensile strength retention 90%, 102%, and 73%, resp., and elongation retention 95%, 95%, and 103%, resp.

- IC ICM C08L077-00
 - ICS C08L067-02; C08L009-02; C08L033-06; C08K003-22; C08L009-00; C08J007-18
- CC 42-11 (Coatings, Inks, and Related Products) Section cross-reference(s): 39
- thermoplastic elastomer fire resistance; chem resistance thermoplastic elastomer; polyester polyether rubber oil resistance; polyamide polyether rubber blend; nitrile rubber blend chem resistance; elec insulator thermoplastic elastomer; alumina fireproofing agent thermoplastic elastomer
- IT **Polyethers**, uses and miscellaneous

(polyamide-, block, rubber, thermoplastic, nitrile rubber blends, contg. alumina trihydrate, fire- and chem. resistant)

- IT Rubber, synthetic
 - (polyamide-polyether, block, thermoplastic, nitrile rubber blends, contg. alumina trihydrate, fire- and chem. resistant)
- IT Polyethers, uses and miscellaneous (polyester-, block, rubber, thermoplastic, nitrile rubber blends, contg. alumina trihydrate, fire- and chem. resistant)
- IT Polyamides, uses and miscellaneous

Polyesters, uses and miscellaneous
 (polyether-, block, rubber, thermoplastic, nitrile
 rubber blends, contg. alumina trihydrate, fire- and chem.
 resistant)